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(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BRAIN

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human brain is described. Also described are single exon nucleic acid probes expressed in the brain and their use in methods for detecting gene expression.

HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BRAIN

CROSS REFERENCE TO RELATED APPLICATIONS

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The present application is a continuation-in-part of U.S. patent application serial nos. 09/632,366, filed August 3, 2000 and 09/608,408, filed June 30, 2000; claims the benefit under 35 U.S.C. s 119(e) of U.S.provisional patent application serial nos. 60/236,359, filed September 27, 2000, 60/234,687, filed September 21, 2000, 60/207,456, filed May 26, 2000, and 60/180,312, filed February 4, 2000; and further claims the benefit under 35 U.S.C. s 119(a) of UK patent application no. 0024263.6, filed October 4, 2000, the disclosures of which are incorporated herein by reference in their entireties.

REFERENCE TO SEQUENCE LISTING AND INCORPORATION BY REFERENCE THEREOF

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The present application includes a Sequence Listing in electronic format, filed pursuant to PCT Administrative Instructions 801 - 806 on a single CD-R disc, in triplicate, containing a file named pto_BRAIN.txt, created 24 January 2001, having 25,840,972 bytes. The Sequence Listing contained in said file on said disc is incorporated herein by reference in its entirety.

Field of the Invention

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The present invention relates to genome-derived single exon microarrays useful for verifying the expression of regions of genomic DNA predicted to encode protein. In particular, the present invention relates to unique genome-derived single exon nucleic acid probes expressed in human

brain and single exon nucleic acid microarrays that include such probes.

Background of the Invention

For almost two decades following the invention of general techniques for nucleic acid sequencing, Sanger et al., Proc. Natl. Acad. Sci. USA 70(4):1209-13 (1973); Gilbert et al., Proc. Natl. Acad. Sci. USA 70(12):3581-4 (1973), these techniques were used principally as tools to further the understanding of proteins — known or suspected — about which a basic foundation of biological knowledge had already been built. In many cases, the cloning effort that preceded sequence identification had been both informed and directed by that antecedent biological understanding.

For example, the cloning of the T cell receptor for antigen was predicated upon its known or suspected cell type-specific expression, by its suspected membrane association, and by the predicted assembly of its gene via T cell-specific somatic recombination. Subsequent sequencing efforts at once confirmed and extended understanding of this family of proteins. Hedrick et al., Nature 308(5955):153-8 (1984).

More recently, however, the development of high
throughput sequencing methods and devices, in concert with
large public and private undertakings to sequence the human
and other genomes, has altered this investigational
paradigm: today, sequence information often precedes
understanding of the basic biology of the encoded protein
product.

One of the approaches to large-scale sequencing is predicated upon the proposition that expressed sequences — that is, those accessible through isolation of mRNA — are of greatest initial interest. This "expressed sequence tag" ("EST") approach has already yielded vast

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amounts of sequence data (see for example Adams et al., Science 252:1651 (1991); Williamson, Drug Discov. Today 4:115 (1999)). For nucleic acids sequenced by this approach, often the only biological information that is known a priori with any certainty is the likelihood of biologic expression itself. By virtue of the species and tissue from which the mRNA had originally been obtained, most such sequences are also annotated with the identity of the species and at least one tissue in which expression appears likely.

More recently, the pace of genomic sequencing has accelerated dramatically. When genomic DNA serves as the initial substrate for sequencing efforts, expression cannot be presumed; often the only a priori biological information about the sequence includes the species and chromosome (and perhaps chromosomal map location) of origin.

With the ever-accelerating pace of sequence accumulation by directed, EST, and genomic sequencing approaches — and in particular, with the accumulation of sequence information from multiple genera, from multiple species within genera, and from multiple individuals within a species — there is an increasing need for methods that rapidly and effectively permit the functions of nucleic sequences to be elucidated. And as such functional information accumulates, there is a further need for methods of storing such functional information in meaningful and useful relationship to the sequence itself; that is, there is an increasing need for means and apparatus for annotating raw sequence data with known or predicted functional information.

Although the increase in the pace of genomic sequencing is due in large part to technological changes in sequencing strategies and instrumentation, Service, Science 280:995 (1998); Pennisi, Science 283: 1822-1823 (1999), there is an important functional motivation as well.

While it was understood that the EST approach would rarely be able to yield sequence information about the noncoding portions of the genome, it now also appears the EST approach is capable of capturing only a fraction of a genome's actual expression complexity.

For example, when the C. elegans genome was fully sequenced, gene prediction algorithms identified over 19,000 potential genes, of which only 7,000 had been found by EST sequencing. C. elegans Sequencing Consortium, 10 Science 282:2012 (1998). Analogously, the recently completed sequence of chromosome 2 of Arabidopsis predicts over 4000 genes, Lin et al., Nature, 402:761 (1999), of which only about 6% had previously been identified via EST sequencing efforts. Although the human genome has the 15 greatest depth of EST coverage, it is still woefully short of surrendering all of its genes. One recent estimate suggests that the human genome contains more than 146,000 genes, which would at this point leave greater than half of the genes undiscovered. It is now predicted that many 20 genes, perhaps 20 to 50%, will only be found by genomic sequencing.

There is, therefore, a need for methods that permit the functional regions of genomic sequence — and most importantly, but not exclusively, regions that

25 function to encode genes — to be identified.

Much of the coding sequence of the human genome is not homologous to known genes, making detection of open reading frames ("ORFs") and predictions of gene function difficult. Computational methods exist for predicting coding regions in eukaryotic genomes. Gene prediction programs such as GRAIL and GRAIL II, Uberbacher et al., Proc. Natl. Acad. Sci. USA 88(24):11261-5 (1991); Xu et al., Genet. Eng. 16:241-53 (1994); Uberbacher et al., Methods Enzymol. 266:259-81 (1996); GENEFINDER, Solovyev et al., Nucl. Acids. Res. 22:5156-63 (1994); Solovyev et al.,

Ismb 5:294-302 (1997); and GENESCAN, Burge et al., J. Mol.
Biol. 268:78-94 (1997), predict many putative genes without
known homology or function. Such programs are known,
however, to give high false positive rates. Burset et al.,
Genomics 34:353-367 (1996). Using a consensus obtained by
a plurality of such programs is known to increase the
reliability of calling exons from genomic sequence.
Ansari-Lari et al., Genome Res. 8(1):29-40 (1998)

Identification of functional genes from genomic

10 data remains, however, an imperfect art. For example, in
reporting the full sequence of human chromosome 21, the
Chromosome 21 Mapping and Sequencing Consortium reports
that prior bioinformatic estimates of human gene number may
need to be revised substantially downwards. Nature

15 405:311-199 (2000); Reeves, Nature 405:283-284 (2000).

Thus, there is a need for methods and apparatus that permit the functions of the regions identified bioinformatically — and specifically, that permit the expression of regions predicted to encode protein — readily to be confirmed experimentally.

Recently, the development of nucleic acid microarrays has made possible the automated and highly parallel measurement of gene expression. Reviewed in Schena (ed.), DNA Microarrays: A Practical Approach

(Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); Nature Genet. 21(1)(suppl):1 - 60 (1999); Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

It is common for microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, such as those from the I.M.A.G.E. consortium, Lennon et al., Genomics 33(1):151-2 (1996), or from the construction of "problem specific" libraries targeted at a particular biological question, R.S. Thomas

et al., Cancer Res. (in press). Such microarrays by definition can measure expression only of those genes found in EST libraries, and thus have not been useful as probes for genes discovered solely by genomic sequencing.

The utility of using whole genome nucleic acid microarrays to answer certain biological questions has been demonstrated for the yeast Saccharomyces cerevisiae. De Risi et al., Science 278:680 (1997). The vast majority of yeast nuclear genes, approximately 95% however, are single exon genes, i.e., lack introns, Lopez et al., RNA 5:1135-1137 (1999); Goffeau et al., Science 274:563-67 (1996), permitting coding regions more readily to be identified. Whole genome nucleic acid microarrays have not generally been used to probe gene expression from more complex eukaryotic genomes, and in particular from those averaging more than one intron per gene.

Diseases of the brain and nervous system are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that

20 contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the brain and nervous system, for the most part these disorders are believed to have polygenic etiologies.

25 There is a need for methods and apparatus that permit prediction, diagnosis and prognosis of diseases of the brain and nervous system particularly those diseases with polygenic etiology.

30 Summary of the Invention

The present invention solves these and other problems in the art by providing methods and apparatus for predicting, confirming, and displaying functional information derived from genomic sequence. The present

invention also provides apparatus for verifying the expression of putative genes identified within genomic sequence.

In particular, the invention provides novel

genome-derived single exon nucleic acid microarrays useful
for verifying the expression of putative genes identified
within genomic sequence.

The present invention also provides compositions and kits for the ready production of nucleic acids

identical in sequence to, or substantially identical in sequence to, probes on the genome-derived single exon microarrays of the present invention.

Accordingly, in a first aspect of the invention, there is provided a spatially-addressable set of single

15 exon nucleic acid probes for measuring gene expression in a sample derived from human brain, comprising a plurality of single exon nucleic acid probes according to any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 12,821 or a complementary sequence, or a portion of such a sequence.

By plurality is meant at least two, suitably at least 20, most suitably at least 100, preferably at least 1000 and, most preferably, upto 5000.

In one embodiment of the first aspect, each of said plurality of probes is separately and addressably amplifiable.

In an alternative embodiment, each of said plurality of probes is separately and addressably isolatable from said plurality.

In a preferred embodiment, each of said plurality of probes is amplifiable using at least one common primer. Preferably, each of said plurality of probes is amplifiable using a first and a second common primer.

In yet another embodiment, said set of single exon nucleic acid probes comprises between 50 - 20,000 probes, for example, 50 - 5000.

Suitably, said set of single exon nucleic acid probes comprises at least 50 - 1000 discrete single exon nucleic acid probes having a sequence as set out in any of SEQ ID NOS.: 1 - 25,434 or a complimentary sequence, or a portion of such a sequence.

Preferably, the average length of the single exon nucleic acid probes is between 200 and 500 bp. It is preferred that the average length should be at least 200bp, suitably at least 250bp, most suitably at least 300bp, preferably at least 400bp and, most preferably, 500 bp.

In another embodiment, the single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.

In another preferred embodiment, said single exon nucleic acid lack homopolymeric stretches of A or T. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.

25 Preferably, a spatially-addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention is is addressably disposed upon a substrate.

Suitable substrates include a filter membrane
30 which may, preferably, be nitrocellulose or nylon. The
13 nylon may preferably, be positively-charged. Other suitable
14 substrates include glass, amorphous silicon, crystalline
15 silicon, and plastic. Further suitable materials include
16 polymethylacrylic, polyethylene, polypropylene,

35 polyacrylate, polymethylmethacrylate, polyvinylchloride,

polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, and mixtures thereof.

In a second aspect of the invention, there is

provided a microarray comprising a spatially addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention.

In one embodiment, a genome-derived single-exon microarray is packaged together with such an ordered set of amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In another aspect, the invention provides genomederived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray. In particular embodiments of this aspect, the present invention provides human single-exon probes that include specifically-hybridizable fragments of SEQ ID Nos. 12,822 - 25,434, wherein the fragment hybridizes at high stringency to an expressed human gene. In particular embodiments, the invention provides single exon probes comprising SEQ ID Nos. 1 - 12,821.

Accordingly, in a third aspect of the invention, there is provided a single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain which is a nucleic acid molecule comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 - 12,821 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human brain.

In one embodiment, a single exon nucleic acid probe in accordance with the third aspect comprises a nucleotide sequence as set out in any of SEQ ID NOs.:

12,822 - 25,434 or a complementary sequence or a fragment thereof.

In a fourth aspect of the invention, there is provided a single exon nucleic acid probe for measuring

5 human gene expression in a sample derived from human brain which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 25,435 ÷ 37,811or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human brain.

Preferably, a single exon nucleic acid probe in accordance with the third or fourth aspects of the invention comprises between at least 15 and 50 contiguous nucleotides of said SEQ ID NO:. It is preferred that the single exon nucleic acid probe comprises at least 15, suitably at least 20, more suitably at least 25 or preferably at least 50 contiguous nucleotides of said SEQ ID NO:.

In another preferred embodiment, a single exon
nucleic acid probe in accordance with the third or fourth
aspects of the invention is between 3kb and 25kb in length.
It is preferred that said probe is no more than 3kb,
suitably no more than 5kb, more suitably no more than 10kb,
preferably 15kb, more preferably 20kb or, most preferably,
no more than 20kb in length.

Preferably, a single exon nucleic acid probe in accordance with either the fifth or sixth aspect of the invention is DNA, preferably single-stranded DNA, RNA or PNA.

In another embodiment of either the third or fourth aspect of the invention, a single exon nucleic acid probe is detectably labeled. Suitable detectable labels include a radionuclide, a fluorescent label or a first member of a specific binding pair. Suitable fluorescent labels include dyes such as cyanine dyes, preferably Cy3

and Cy5 although other suitable dyes will be known to those skilled in the art.

In a particularly preferred embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks prokaryotic and bacteriophage vector sequence. In yet another embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks homopolymeric stretches of A or T.

In a fifth aspect of the invention, there is provided an amplifiable nucleic acid composition, comprising:

the single exon nucleic acid probe in accordance with either of the third or fourth aspects of the invention; and at least one nucleic acid primer;

wherein said at least one primer is sufficient to prime enzymatic amplification of said probe.

In an sixth aspect of the invention, there is provided a method of measuring gene expression in a sample derived from human brain, comprising:

contacting the single exon microarray in accordance with the second aspect of the invention, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human brain; and then

measuring the label detectably bound to each probe of said microarray.

In a seventh aspect of the invention, there is provided a method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then

detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are

derived from mRNA from the brain of said eukaryote, said probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a single exon 5 microarray in accordance with the first aspect of the invention, and said fragment is selectively hybridizable at high stringency.

In a eighth aspect of the invention, there is provided a method of assigning exons to a single gene,

10 comprising:

identifying a plurality of exons from genomic sequence in accordance with the seventh aspect of the invention; and then

measuring the expression of each of said exons in

15 a plurality of tissues and/or cell types using
hybridization to single exon microarrays having a probe
with said exon,

wherein a common pattern of expression of said exons in said plurality of tissues and/or cell types 20 indicates that the exons should be assigned to a single gene.

In an ninth aspect of the invention, there is provided a nucleic acid sequence as set out in any of SEQ ID NOs: 1 - 25,434 wherein said sequence encodes a peptide.

In a tenth aspect of the invention, there is provided a peptide encoded by a sequence comprising a sequence as set out in any of SEQ ID NOs: 12,822 - 25,434, or a complementary sequence or coding portion thereof.

In a preferred embodiment, a peptide may be
30 encoded by a sequence comprising a sequence set out in any
of SEQ ID NOS.: 1 -12,821.

In a further aspect, the invention provides peptides comprising an amino acid sequence translated from the DNA fragments, said amino acid sequences comprising SEQ ID NOS.: 25,435 - 37,811.

Accordingly in a eleventh aspect of the invention there is provided a peptide comprising a sequence as set out in any of SEQ ID NOs: 25,435-37,811, or fragment thereof.

In another aspect, the invention provides means for displaying annotated sequence, and in particular, for displaying sequence annotated according to the methods and apparatus of the present invention. Further, such display can be used as a preferred graphical user interface for electronic search, query, and analysis of such annotated sequence.

Detailed Description of the Invention

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Definitions

As used herein, the term "microarray" and phrase "nucleic acid microarray" refer to a substrate-bound collection of plural nucleic acids, hybridization to each of the plurality of bound nucleic acids being separately detectable. The substrate can be solid or porous, planar or non-planar, unitary or distributed.

As so defined, the term "microarray" and phrase "nucleic acid microarray" include all the devices so called in Schena (ed.), DNA Microarrays: A Practical Approach (Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); Nature Genet. 21(1)(suppl):1 - 60 (1999); and Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376). As so defined, the term "microarray" and phrase "nucleic acid microarray" further include substrate-bound collections of plural nucleic acids in which the nucleic acids are distributably disposed on a plurality of beads, rather than on a unitary planar substrate, as is described, inter alia, in Brenner

et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); in such case, the term "microarray" and phrase "nucleic acid microarray" refer to the plurality of beads in aggregate.

As used herein with respect to a nucleic acid microarray, the term "probe" refers to the nucleic acid that is, or is intended to be, bound to the substrate; in such context, the term "target" thus refers to nucleic acid intended to be bound thereto by Watson-Crick

10 complementarity. As used herein with respect to solution phase hybridization, the term "probe" refers to the nucleic acid of known sequence that is detectably labeled.

As used herein, the expression "probe comprising SEQ ID NO.", and variants thereof, intends a nucleic acid probe, at least a portion of which probe has either (i) the sequence directly as given in the referenced SEQ ID NO., or (ii) a sequence complementary to the sequence as given in the referenced SEQ ID NO., the choice as between sequence directly as given and complement thereof dictated by the requirement that the probe hybridize to mRNA.

As used herein, the term "open reading frame" and the equivalent acronym "ORF" refer to that portion of an exon that can be translated in its entirety into a sequence of contiguous amino acids i.e. a nucleic acid sequence

25 that, in at least one reading frame, does not possess stop codons; the term does not require that the ORF encode the entirety of a natural protein.

As used herein, the term "amplicon" refers to a PCR product amplified from human genomic DNA, containing the predicted exon.

As used herein the term "exon" refers to the consensus prediction of the various exon and gene predicting algorithms i.e. a nucleic acid sequence bioinformatically predicted to encode a portion of a natural protein.

As used herein, the term "peptide" refers to a sequence of amino acids. The sequences referred to as PEPTIDE SEQ ID NOS.: are the predicted peptide sequences that would be translated from one of the exons, or a portion thereof set out in exon SEQ ID NOS.:. The codons encoding the peptide are wholly contained within the exon.

As used herein, a "portions" of a defined nucleotide sequence or sequences can be and, preferably, are fragments unique to that sequence or to one or a combination of those sequences. A fragment unique to a nucleic acid molecule is one that is a signature for the larger nucleic acid molecule.

As used herein, the phrase "expression of a probe" and its linguistic variants means that the ORF present within the probe, or its complement, is present within a target mRNA.

As used herein, "stringent conditions" refers to parameters well known to those skilled in the art. When a nucleic acid molecule is said to be hybridisable to another of a given sequence under "stringent conditions" it is meant that it is homologous to the given sequence.

As used herein, the phrase "specific binding pair" intends a pair of molecules that bind to one another with high specificity. Binding pairs are said to exhibit specific binding when they exhibit avidity of at least 10⁷, preferably at least 10⁸, more preferably at least 10⁹ liters/mole. Nonlimiting examples of specific binding pairs are: antibody and antigen; biotin and avidin; and biotin and streptavidin.

As used herein with respect to the visual display of annotated genomic sequence, the term "rectangle" means any geometric shape that has at least a first and a second border, wherein the first and second borders each are capable of mapping uniquely to a point of another visual object of the display.

As used herein, a "Mondrian" means a visual display in which a single genomic sequence is annotated with predicted and experimentally confirmed functional information.

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Brief Description of the Drawings

The present invention is further illustrated with reference to the following non-limiting figures and examples in which:

FIG. 1 illustrates a process for predicting functional regions from genomic sequence, confirming the functional activity of such regions experimentally, and associating and displaying the data so obtained in meaningful and useful relationship to the original sequence data;

FIG. 2 further elaborates that portion of the process schematized in FIG. 1 for predicting functional regions from genomic sequence;

FIG. 3 illustrates a Mondrian visual display;
FIG. 4 presents a Mondrian showing a hypothetical annotated genomic sequence;

FIG. 5 is a histogram showing the distribution of ORF length and PCR products as obtained, with ORF length shown in black and PCR product length shown in dotted lines;

FIG. 6 is a histogram showing the distribution, among exons predicted according to the methods described,

30 of expression as measured using simultaneous two color hybridization to a genome-derived single exon microarray. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), or expressed in all tissues tested ("10");

FIG. 7 is a pictorial representation of the expression of verified sequences that showed expression with signal intensity greater than 3 in at least one tissue, with: FIG. 7A showing the expression as measured by 5 microarray hybridization in each of the 10 measured tissues, and the expression as measured "bioinformatically" by query of EST, NR and SwissProt databases; with FIG. 7B showing the legend for display of physical expression (ratio) in FIG. 7A; and with FIG. 7C showing the legend for scoring EST hits as depicted in FIG. 7A;

FIG. 8 shows a comparison of normalized CY3 signal intensity for arrayed sequences that were identical to sequences in existing EST, NR and SwissProt databases or that were dissimilar (unknown), where black denotes the 15 signal intensity for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 (1 x 10^{-30}) ("unknown") and a dotted line denotes sequence-verified spots with a BLAST expect ("E") value of less than 1e-30 (1 $\times 10^{-30}$) ("known");

FIG. 9 presents a Mondrian of BAC AC008172 (bases 20 25,000 to 130,000), containing the carbamyl phosphate synthetase gene (AF154830.1); and

FIG. 10 is a Mondrian of BAC A049839.

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Methods and Apparatus for Predicting, Confirming, Annotating, and Displaying Functional Regions From Genomic Sequence Data

FIG. 1 is a flow chart illustrating in broad 30 outline a process for predicting functional regions from genomic sequence, confirming and characterizing the functional activity of such regions experimentally, and then associating and displaying the information so obtained 35 in meaningful and useful relationship to the original

sequence data.

The initial input into process 10 of the present invention is drawn from one or more databases 100 containing genomic sequence data. Because genomic sequence is usually obtained from subgenomic fragments, the sequence data typically will be stored in a series of records corresponding to these subgenomic sequenced fragments. Some fragments will have been catenated to form larger contiguous sequences ("contigs"); others will not. A finite percentage of sequence data in the database will typically be erroneous, consisting inter alia of vector sequence, sequence created from aberrant cloning events, sequence of artificial polylinkers, and sequence that was erroneously read.

Each sequence record in database 100 will minimally contain as annotation a unique sequence identifier (accession number), and will typically be annotated further to identify the date of accession, species of origin, and depositor. Because database 100 can contain nongenomic sequence, each sequence will typically be annotated further to permit query for genomic sequence. Chromosomal origin, optionally with map location, can also be present. Data can be, and over time increasingly will be, further annotated with additional information, in part through use of the present invention, as described below. Annotation can be present within the data records, in information external to database 100 and linked to the records thereto, or through a combination of the two.

Databases useful as genomic sequence database 100
in the present invention include GenBank, and particularly include several divisions thereof, including the htgs(draft), NT (nucleotide, command line), and NR (nonredundant) divisions. GenBank is produced by the National Institutes of Health and is maintained by the National Center for Biotechnology Information (NCBI).

Databases of genomic sequence from species other than human, such as mouse, rat, Arabidopsis, C. elegans, C. brigsii, Drosophila, zebra fish, and other higher eukaryotic organisms will also prove useful as genomic sequence database 100.

Genomic sequence obtained by query of genomic sequence database 100 is then input into one or more processes 200 for identification of regions therein that are predicted to have a biological function as specified by the user. Such functions include, but are not limited to, encoding protein, regulating transcription, regulating message transport after transcription into mRNA, regulating message splicing after transcription into mRNA, of regulating message degradation after transcription into 15 mRNA, and the like. Other functions include directing somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, and the like.

The particular genomic sequence to be input into process 200 will depend upon the function for which relevant sequence is to be identified as well as upon the approach chosen for such identification. Process step 200 can be iterated to identify different functions within a given genomic region. In such case, the input often will be different for the several iterations.

Sequences predicted to have the requisite function by process 200 are then input into process 300, where a subset of the input sequences suitable for experimental confirmation is identified. Experimental confirmation can involve physical and/or bioinformatic assay. Where the subsequent experimental assay is bioinformatic, rather than physical, there are fewer constraints on the sequences that can be tested, and in this latter case therefore process 300 can output the entirety of the input sequence.

The subset of sequences output from process 300 is then used in process 400 for experimental verification and characterization of the function predicted in process 200, which experimental verification can, and often will, include both physical and bioinformatic assay.

Process 500 annotates the sequence data with the functional information obtained in the physical and/or bioinformatic assays of process 400. Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the sequence data record itself, by linking records in a hierarchical or relational database, by linking to external databases, by a combination thereof, or by other means well known within the database arts. The data can even be submitted for incorporation into databases maintained by others, such as GenBank, which is maintained by NCBI.

As further noted in FIG. 1, additional annotation can be input into process 500 from external sources 600.

The annotated data is then displayed in process 800, either before, concomitantly with, or after optional storage 700 on nontransient media, such as magnetic disk, optical disc, magnetooptical disk, flash memory, or the like.

25 FIG. 1 shows that the experimental data output from process 400 can be used in each preceding step of process 10: e.g., facilitating identification of functional sequences in process 200, facilitating identification of an experimentally suitable subset thereof in process 300, and 30 facilitating creation of physical and/or informational substrates for, and performance of subsequent assay, of functional sequences in process 400.

Information from each step can be passed directly to the succeeding process, or stored in permanent or interim form prior to passage to the succeeding process.

Often, data will be stored after each, or at least a plurality, of such process steps. Any or all process steps can be automated.

FIG. 2 further elaborates the prediction of functional sequence within genomic sequence according to process 200.

Genomic sequence database 100 is first queried 20 for genomic sequence.

The sequence required to be returned by query 20 10 will depend, in the first instance, upon the function to be identified.

For example, genomic sequences that function to encode protein can be identified *inter alia* using gene prediction approaches, comparative sequence analysis

- approaches, or combinations of the two. In gene prediction analysis, sequence from one genome is input into process 200 where at least one, preferably a plurality, of algorithmic methods are applied to identify putative coding regions. In comparative sequence analysis, by contrast,
- corresponding, e.g., syntenic, sequence from a plurality of sources, typically a plurality of species, is input into process 200, where at least one, possibly a plurality, of algorithmic methods are applied to compare the sequences and identify regions of least variability.
- 25 The exact content of query 20 will also depend upon the database queried. For example, if the database contains both genomic and nongenomic sequence, perhaps derived from multiple species, and the function to be determined is protein coding regions in human genomic sequence, the query will accordingly require that the sequence returned be genomic and derived from humans.

Query 20 can also incorporate criteria that compel return of sequence that meets operative requirements of the subsequent analytical method. Alternatively, or in addition, such operative criteria can be enforced in

subsequent preprocess step 24.

For example, if the function sought to be identified is protein coding, query 20 can incorporate criteria that return from genomic sequence database 100 only those sequences present within contigs sufficiently long as to have obviated substantial fragmentation of any given exon among a plurality of separate sequence fragments.

Such criteria can, for example, consist of a

required minimal individual genomic sequence fragment
length, such as 10 kb, more typically 20 kb, 30 kb, 40kb,
and preferably 50 kb or more, as well as an optional
further or alternative requirement that sequence from any
given clone, such as a bacterial artificial chromosome

("BAC"), be presented in no more than a finite maximal
number of fragments, such as no more than 20 separate
pieces, more typically no more than 15 fragments, even more
typically no more than about 10 - 12 fragments.

Results using the present invention have shown
that genomic sequence from bacterial artificial chromosomes
(BACs) is sufficient for gene prediction analysis according
to the present invention if the sequence is at least 50 kb
in length, and if additionally the sequence from any given
BAC is presented in fewer than 15, and preferably fewer
than 10, fragments. Accordingly, query 20 can incorporate
a requirement that data accessioned from BAC sequencing be
in fewer than 15, preferably fewer than 10, fragments.

An additional criterion that can be incorporated into the query can be the date, or range of dates, of

30 sequence accession. Although the process has been described above as if genomic sequence database 100 were static, it is of course understood that the genomic sequence databases need not be static, and indeed are typically updated on a frequent, even hourly, basis. Thus, as further described in Examples 1 and 2, infra, it is

possible to query the database for newly added sequence, either newly added after an absolute date, or newly added relative to a prior analysis performed using the methods and apparatus of the present invention. In this way, the process herein described can incorporate a dynamic, temporal component.

One utility of such temporal limitation is to identify, from newly accessioned genomic sequence, the presence of novel genes, particularly those not previously identified by EST sequencing (or other sequencing efforts that are similarly based upon gene expression). As further described in Example 1, such an approach has shown that newly accessioned human genomic sequence, when analyzed for sequences that function to encode protein, readily identifies genes that are novel over those in existing EST and other expression databases. This makes the methods of the present invention extremely powerful gene discovery tools. And as would be appreciated, such gene discovery can be performed using genomic sequence from species other than human.

If query 20 incorporates multiple criteria, such as above-described, the multiple criteria can be performed as a series of separate queries or as a single query, depending in part upon the query language, the complexity of the query, and other considerations well known in the database arts.

If query 20 returns no genomic sequence meeting the query criteria, the negative result can be reported by process 22, and process 200 (and indeed, entire process 10) ended 23, as shown. Alternatively, or in addition to report and termination of the initial inquiry, a new query 20 can be generated that takes into account the initial negative result.

When query 20 returns sequence meeting the query 35 criteria, the returned sequence is then passed to optional

preprocessing 24, suitable and specific for the desired analytical approach and the particular analytical methods thereof to be used in process 25.

Preprocessing 24 can include processes suitable for many approaches and methods thereof, as well as processes specifically suited for the intended subsequent analysis.

Preprocessing 24 suitable for most approaches and methods will include elimination of sequence irrelevant to, or that would interfere with, the subsequent analysis. Such sequence includes repetitive sequence, such as Alu repeats and LINE elements, vector sequence, artificial sequence, such as artificial polylinkers, and the like. Such removal can readily be performed by identification and subsequent masking of the undesired sequence.

Identification can be effected by comparing the genomic sequence returned by query 20 with public or private databases containing known repetitive sequence, vector sequence, artificial sequence, and other artifactual sequence. Such comparison can readily be done using programs well known in the art, such as CROSS_MATCH, or by proprietary sequence comparison programs the engineering of which is well within the skill in the art.

Alternatively, or in addition, undesirable,
including artifactual, sequence can be identified
algorithmically without comparison to external databases
and thereafter removed. For example, synthetic polylinker
sequence can be identified by an algorithm that identifies
a significantly higher than average density of known
restriction sites. As another example, vector sequence can
be identified by algorithms that identify nucleotide or
codon usage at variance with that of the bulk of the
genomic sequence.

Once identified, undesired sequence can be removed. Removal can usefully be done by masking the

undesired sequence as, for example, by converting the specific nucleotide references to one that is unrecognized by the subsequent bioinformatic algorithms, such as "X". Alternatively, but at present less preferred, the undesired sequence can be excised from the returned genomic sequence, leaving gaps.

Preprocessing 24 can further include selection from among duplicative sequences of that one sequence of highest quality. Higher quality can be measured as a lower percentage of, fewest number of, or least densely clustered occurrence of ambiguous nucleotides, defined as those nucleotides that are identified in the genomic sequence using symbols indicating ambiguity. Higher quality can also or alternatively be valued by presence in the longest contig.

Preprocessing 24 can, and often will, also include formatting of the data as specifically appropriate for passage to the analytical algorithms of process 25. Such formatting can and typically will include, inter alia, addition of a unique sequence identifier, either derived from the original accession number in genomic sequence database 100, or newly applied, and can further include additional annotation. Formatting can include conversion from one to another sequence listing standard, such as conversion to or from FASTA or the like, depending upon the input expected by the subsequent process.

Preprocessing, which can be optional depending upon the function desired to be identified and the informational requirements of the methods for effecting such identification, is followed by sequence processing 25, where sequences with the desired function are identified within the genomic sequence.

As mentioned above, such functions can include, but are not limited to, encoding protein, regulating

35 transcription, regulating message transport after

transcription into mRNA, regulating message splicing after transcription, of regulating message degradation, and the like. Other functions include directing somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, or the like.

The methods of the present invention are particularly useful for gene discovery, that is, for identifying, from genomic sequence, regions that function to encode genes, and in a particularly useful embodiment, for identifying regions that function to encode genes not hitherto identified by expression-based or directed cloning and sequencing. In conjunction with verification using the novel single exon microarrays of the present invention, as further described below, the methods herein described become powerful gene discovery tools.

Accordingly, in a preferred embodiment of the present invention, process 25 is used to identify putative coding regions. Two preferred approaches in process 25 for identifying sequence that encodes putative genes are gene prediction and comparative sequence analysis.

Gene prediction can be performed using any of a number of algorithmic methods, embodied in one or more software programs, that identify open reading frames (ORFs) using a variety of heuristics, such as GRAIL, DICTION, and GENEFINDER. Comparative sequence analysis similarly can be performed using any of a variety of known programs that identify regions with lower sequence variability.

As further described in Example 1, below, gene

finding software programs yield a range of results. For
the newly accessioned human genomic sequence input in
Example 1, for example, GRAIL identified the greatest
percentage of genomic sequence as putative coding region,
2% of the data analyzed; GENEFINDER was second, calling 1%;
and DICTION yielded the least putative coding region, with

0.8% of genomic sequence called as coding region.

Increased reliability can be obtained when consensus is required among several such methods. Although discussed herein particularly with respect to exon calling, consensus among methods will in general increase reliability of predicting other functions as well.

Thus, as indicated by query 26, sequence processing 25, optionally with preprocessing 24, can be repeated with a different method, with consensus among such iterations determined and reported in process 27.

Process 27 compares the several outputs for a given input genomic sequence and identifies consensus among the separately reported results. The consensus itself, as well as the sequence meeting that consensus, is then stored in process 29a, displayed in process 29b, and/or output to process 300 for subsequent identification of a subset thereof suitable for assay.

Multiple levels of consensus can be calculated and reported by process 27. For example, as further described in Example 1, infra, process 27 can report consensus as between all specific pairs of methods of gene prediction, as consensus among any one or more of the pairs of methods of gene prediction, or as among all of the gene prediction algorithms used. Thus, in Example 1, process 27 reported that GRAIL and GENEFINDER programs agreed on 0.7% of genomic sequence, that GRAIL and DICTION agreed on 0.5% of genomic sequence, and that the three programs together agreed on 0.25% of the data analyzed. Put another way, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding region.

Furthermore, consensus can be required among different approaches to identifying a chosen function.

For example, if the function desired to be identified is coding of protein sequence, and a first used approach to exon calling is gene prediction, the process

can be repeated on the same input sequence, or subset thereof, with another approach, such as comparative sequence analysis. In such a case, where comparative sequence analysis follows gene prediction, the comparison can be performed not only on genomic nucleic acid sequence, but additionally or alternatively can be performed on the predicted amino acid sequence translated from the ORFs prior identified by the gene prediction approach.

Although shown as an iterative process, the

10 multiple analyses required to achieve consensus can be done
in series, in parallel, or some combination thereof.

Predicted functional sequence, optionally representing a consensus among a plurality of methods and approaches for determination thereof, is passed to process 300 for identification of a subset thereof for functional assay.

In the preferred embodiment of the methods of the present invention, wherein the function sought to be identified is protein coding, process 300 is used to 20 identify a subset thereof suitable for experimental verification by physical and/or bioinformatic approaches.

For example, putative ORFs identified in process 200 can be classified, or binned, bioinformatically into putative genes. This binning can be based inter alia upon consideration of the average number of exons/gene in the species chosen for analysis, upon density of exons that have been called on the genomic sequence, and other empirical rules. Thereafter, one or more among the genespecific ORFs can be chosen for subsequent use in gene expression assay.

Where such subsequent gene expression assay uses amplified nucleic acid, considerations such as desired amplicon length, primer synthesis requirements, putative exon length, sequence GC content, existence of possible secondary structure, and the like can be used to identify

and select those ORFs that appear most likely successfully to amplify. Where subsequent gene expression assay relies upon nucleic acid hybridization, whether or not using amplified product, further considerations involving

5 hybridization stringency can be applied to identify that subset of sequences that will most readily permit sequence-specific discrimination at a chosen hybridization and wash stringency. One particular such consideration is avoidance of putative exons that span repetitive sequence; such sequence can hybridize spuriously to nonspecific message, reducing specific signal in the hybridization.

For bioinformatic assay, there are fewer constraints on the sequences that can be tested experimentally, and in this latter case therefore process 300 can output the entirety of the input sequence.

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The subset of sequences identified by process 300 as suitable for use in assay is then used in process 400 to create the physical and/or informational substrate for experimental verification of the predictions made in process 200, and thereafter to assay those substrates.

As mentioned, the methods of the present invention are particularly useful for identifying potential coding regions within genomic sequence. In a preferred embodiment of process 400, therefore, the expression of the sequences predicted to encode protein is verified. The combination of the predictive and experimental methods provides a powerful gene discovery engine.

Thus, in another aspect, the present invention provides methods and apparatus for verifying the expression of putative genes identified within genomic sequence. In particular, the invention provides a novel method of verifying gene expression in which expression of predicted ORFs is measured and confirmed using a novel type of nucleic acid microarray, the genome-derived single exon nucleic acid microarrays of the present invention.

Putative ORFs as predicted by a consensus of gene calling, particularly gene prediction, algorithms in process 200, and as further identified as suitable by process 300, are amplified from genomic DNA using the 5 polymerase chain reaction (PCR). Although PCR is conveniently used, other amplification approaches can also be used.

Amplification schemes can be designed to capture the entirety of each predicted ORF in an amplicon with 10 minimal additional (that is, intronic or intergenic) sequence. Because ORFs predicted from human genomic sequence using the methods of the present invention differ in length, such an approach results in amplicons of varying length.

15 However, most predicted ORFs are shorter than 500 bp in length, and although amplicons of at least about 100 or 200 base pairs can be immobilized as probes on nucleic acid microarrays, early experimental results using the methods of the present invention have suggested that longer 20 amplicons, at least about 400 or 500 base pairs, are more effective. Furthermore, certain advantages derive from application to the microarray of amplicons of defined size.

Therefore, amplification schemes can alternatively, and preferably, be designed to amplify 25 regions of defined size, preferably at least about 300, 400 or 500 bp, centered about each predicted ORF. Such an approach results in a population of amplicons of limited size diversity, but that typically contain intronic and/or intergenic nucleic acid in addition to putative ORF.

Conversely, somewhat fewer than 10% of ORFs predicted from human genomic sequence according to the methods of the present invention exceed 500 bp in length. Portions of such extended ORFs, preferably at least about 300,400 or 500 bp in length, can be amplified. However, it 35 has been discovered that the percentage success at

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amplifying pieces of such ORFs is low, and that such putative exons are more effectively amplified when larger fragments, at least about 1000 or 1500 bp, and even as large as 2000 bp are amplified.

thus input into one or more primer design programs, such as PRIMER3 (available online for use at http://www-genome.wi.mit.edu/cgi-bin/primer/), with a goal of amplifying at least about 500 base pairs of genomic
sequence centered within or about ORFs predicted to be no more than about 500 bp, or at least about 1000 - 1500 bp of genomic sequence for ORFs predicted to exceed 500 bp in length, and the primers synthesized by standard techniques. Primers with the requisite sequences can be purchased
commercially or synthesized by standard techniques.

Conveniently, a first predetermined sequence can be added commonly to the ORF-specific 5' primer and a second, typically different, predetermined sequence commonly added to each 3' ORF-unique primer. This serves to immortalize the amplicon, that is, serves to permit further amplification of any amplicon using a single set of primers complementary respectively to the common 5' and common 3' sequence elements. The presence of these "universal" priming sequences further facilitates later sequence verification, providing a sequence common to all amplicons at which to prime sequencing reactions. The common 5' and 3' sequences further serve to add a cloning site should any of the ORFs warrant further study.

Such predetermined sequence is usefully at least 30 about 10, 12 or 15 nt in length, and usually does not exceed about 25 nt in length. The "universal" priming sequences used in the examples presented *infra* were each 16 nt long.

The genomic DNA to be used as substrate for amplification will come from the eukaryotic species from

which the genomic sequence data had originally been obtained, or a closely related species, and can conveniently be prepared by well known techniques from somatic or germline tissue or cultured cells of the organism. See, e.g., Short Protocols in Molecular Biology: A Compendium of Methods from Current Protocols in Molecular Biology, Ausubel et al. (eds.), 4th edition (April 1999), John Wiley & Sons (ISBN: 047132938X) and Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd edition (December 1989), Cold Spring Harbor Laboratory Press (ISBN: 0879693096). Many such prepared genomic DNAs are available commercially, with the human genomic DNAs additionally having certification of donor informed consent.

15 Although the intronic and intergenic material flanking putative coding regions in the amplicons could potentially interfere with hybridizations during microarray experiments, we have found, surprisingly, that differential expression ratios are not significantly affected. Rather, 20 the predominant effect of exon size is to alter the absolute signal intensity, rather than its ratio. Equally surprising, the art had suggested that single exon probes would not provide sufficient signal intensity for high stringency hybridization analyses; we find that such probes 25 not only provide adequate signal, but have substantial advantages, as herein described.

After partial purification, as by size exclusion spin column, with or without confirmation as to amplicon quality as by gel electrophoresis, each amplicon (single exon probe) is disposed in an array upon a support substrate.

Methods for creating microarrays by deposition and fixation of nucleic acids onto support substrates are well known in the art (Reviewed by Schena et al., see above).

Typically, the support substrate will be glass, although other materials, such as amorphous or crystalline silicon or plastics. Such plastics include polymethylacrylic, polyethylene, polypropylene,

5 polyacrylate, polymethylmethacrylate, polyvinylchloride, polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, or mixtures thereof, can also be used. Typically, the support will be rectangular, although other shapes, particularly circular disks and even spheres, present certain advantages. Particularly advantageous alternatives to glass slides as support substrates for array of nucleic acids are optical discs, as described in WO 98/12559.

The amplified nucleic acids can be attached covalently to a surface of the support substrate or, more typically, applied to a derivatized surface in a chaotropic agent that facilitates denaturation and adherence by presumed noncovalent interactions, or some combination thereof.

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Robotic spotting devices useful for arraying nucleic acids on support substrates can be constructed using public domain specifications (The MGuide, version 2.0, http://cmgm.stanford.edu/pbrown/mguide/index.html), or can conveniently be purchased from commercial sources (MicroArray GenII Spotter and MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Spotting can also be effected by printing methods, including those using ink jet technology.

As is well known in the art, microarrays typically also contain immobilized control nucleic acids. For controls useful in providing measurements of background signal for the genome-derived single exon microarrays of the present invention, a plurality of *E. coli* genes can readily be used. As further described in Example 1, 16 or

32 *E. coli* genes suffice to provide a robust measure of background noise in such microarrays.

As is well known in the art, the amplified product disposed in arrays on a support substrate to create a nucleic acid microarray can consist entirely of natural nucleotides linked by phosphodiester bonds, or alternatively can include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained in the hybridization.

If enzymatic amplification is used to produce the immobilized probes, the amplifying enzyme will impose certain further constraints upon the types of nucleic acid analogs that can be generated.

Although particularly described herein as using
high density microarrays constructed on planar substrates,
the methods of the present invention for confirming the
expression of ORFs predicted from genomic sequence can use
any of the known types of microarrays, as herein defined,
including lower density planar arrays, and microarrays on
nonplanar, nonunitary, distributed substrates.

For example, gene expression can be confirmed using hybridization to lower density arrays, such as those constructed on membranes, such as nitrocellulose, nylon, and positively-charged derivatized nylon membranes.

25 Further, gene expression can also be confirmed using nonplanar, bead-based microarrays such as are described in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); U.S. Patent No. 6,057,107; and U.S. Patent No. 5,736,330. In theory, a packed collection of such beads provides in aggregate a higher density of nucleic acid probe than can be achieved with spotting or lithography techniques on a single planar substrate.

Planar microarrays on solid substrates, however, provide certain useful advantages, including high
throughput and compatibility with existing readers. For

example, each standard microscope slide can include at least 1000, typically at least 2000, preferably 5000 and upto 10,000 - 50,000 or more nucleic acid probes of discrete sequence. The number of sequences deposited will 5 depend on their required application.

Each putative gene can be represented in the array by a single predicted ORF. Alternatively, genes can be represented by more than one predicted ORF. For purposes of measuring differential splicing, more than one 10 predicted ORF will be provided for a putative gene. And as is well known in the art, each probe of defined sequence, representing a single predicted ORF, can be deposited in a plurality of locations on a single microarray to provide redundancy of signal.

The genome-derived single exon microarrays described above differ in several fundamental and advantageous ways from microarrays presently used in the gene expression art, including (1) those created by deposition of mRNA-derived nucleic acids, (2) those created 20 by in situ synthesis of oligonucleotide probes, and (3) those constructed from yeast genomic DNA.

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Most nucleic acid microarrays that are in use for study of eukaryotic gene expression have as immobilized probes nucleic acids that are derived — either directly or 25 indirectly - from expressed message. As discussed above, it is common, for example, for such microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, see Lennon et al., or from the de novo construction of "problem specific" 30 libraries targeted at a particular biological question, R.S. Thomas et al., Cancer Res. (in press). Such microarrays are herein collectively denominated "EST. microarrays".

Such EST microarrays by definition can measure 35 expression only of those genes found in EST libraries,

shown herein to represent only a fraction of expressed genes. Furthermore, such libraries - and thus microarrays based thereupon - are biased by the tissue or cell type of message origin, by the expression levels of the respective 5 genes within the tissues, and by the ability of the message successfully to have been reverse-transcribed and cloned.

Thus, as further discussed in Example 1, the methods of the present invention enable sequences that do not appear in EST or other expression databases to be 10 determined - subsequently arrayed for expression measurements could not, therefore, have been represented as probes on an EST microarray. And as further demonstrated in the examples, infra, the remaining population of genes identified from genomic sequence by the methods of the 15 present invention - that is, the one third of sequences that had previously been accessioned in EST or other expression databases - are biased toward genes with higher expression levels.

Representation of a message in an EST and/or cDNA 20 library depends upon the successful reverse transcription, optionally but typically with subsequent successful cloning, of the message. This introduces substantial bias into the population of probes available for arraying in EST microarrays.

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In contrast, neither reverse transcription nor cloning is required to produce the probes arrayed on the genome-derived single exon microarrays of the present invention. And although the ultimate deposition of a probe on the genome-derived single exon microarray of the present 30 invention depends upon a successful amplification from genomic material, a priori knowledge of the sequence of the desired amplicon affords greater opportunity to recover any given probe sequence recalcitrant to amplification than is afforded by the requirement for successful reverse 35 transcription and cloning of unknown message in EST

approaches.

Thus, the genome-derived single exon microarrays of the present invention present a far greater diversity of probes for measuring gene expression, with far less bias, than do EST microarrays presently used in the art.

As a further consequence of their ultimate origin from expressed message, the probes in EST microarrays often contain poly-A (or complementary poly-T) stretches derived from the poly-A tail of mature mRNA. These homopolymeric stretches contribute to cross-hybridization, that is, to a spurious signal occasioned by hybridization to the homopolymeric tail of a labeled cDNA that lacks sequence homology to the gene-specific portion of the probe.

In contrast, the probes arrayed in the genomederived single exon microarrays of the present invention
lack homopolymeric stretches derived from message
polyadenylation, and thus can provide more specific signal.
Typically, at least about 50, 60 or 75% of the probes on
the genome-derived single exon microarrays of the present
invention lack homopolymeric regions consisting of A or T,
where a homopolymeric region is defined for purposes herein
as stretches of 25 or more, typically 30 or more, identical
nucleotides.

A further distinction, which also affects the

25 specificity of hybridization, is occasioned by the typical
derivation of EST microarray probes from cloned material.
Because much of the probe material disposed as probes on
EST microarrays is excised or amplified from plasmid,
phage, or phagemid vectors, EST microarrays typically

30 include a fair amount of vector sequence, more so when the
probes are amplified, rather than excised, from the vector.

In contrast, the vast majority of probes in the genome-derived single exon microarrays of the present invention contain no prokaryotic or bacteriophage vector sequence, having been amplified directly or indirectly from

genomic DNA. Typically, therefore, at least about 50, 60, 70 or 80% or more of individual exon-including probes disposed on a genome-derived single exon microarray of the present invention lack vector sequence, and particularly lack sequences drawn from plasmids and bacteriophage. Preferably, at least about 85, 90 or more than 90% of exon-including probes in the genome-derived single exon microarray of the present invention lack vector sequence. With attention to removal of vector sequences through preprocessing 24, percentages of vector-free exon-including probes can be as high as 95 - 99%. The substantial absence of vector sequence from the genome-derived single exon microarrays of the present invention results in greater

As a further consequence of excision or amplification of probes from vectors in construction of EST microarrays, the probes arrayed thereon often contain artificial sequence, derived from vector polylinker

20 multiple cloning sites, at both 5' and 3' ends. The probes disposed upon the genome-derived single exon microarrays need have no such artificial sequence appended thereto.

specificity during hybridization, since spurious cross-

15 hybridization to a probe vector sequence is reduced.

As mentioned above, however, the ORF-specific primers used to amplify putative ORFs can include

25 artificial sequences, typically 5' to the ORF-specific primer sequence, useful for "universal" (that is, independent of ORF sequence) priming of subsequent amplification or sequencing reactions. When such "universal" 5' and/or 3' priming sequences are appended to the amplification primers, the probes disposed upon the genome-derived single exon microarray will include artificial sequence similar to that found in EST microarrays. However, the genome-derived single exon microarray of the present invention can be made without such sequences, and if so constructed, presents an even

smaller amount of nonspecific sequence that would contribute to nonspecific hybridization.

Yet another consequence of typical use of cloned material as probes in EST microarrays is that such

5 microarrays contain probes that result from cloning artifacts, such as chimeric molecules containing coding region of two separate genes. Derived from genomic material, typically not thereafter cloned, the probes of the genome-derived single exon microarrays of the present invention lack such cloning artifacts, and thus provide greater specificity of signal in gene expression measurements.

A further consequence of the cloned origin of probes on many EST microarrays is that the individual

15 probes often have disparate sizes, which can cause the optimal hybridization stringency to vary among probes on a single microarray. In contrast, as discussed above, the probes arrayed on the genome-derived single exon microarrays of the present invention can readily be

20 designed to have a narrow distribution in sizes, with the range of probe sizes no greater than about 10% of the average size, typically no greater than about 5% of the average probe size.

Because of their origin from fully- or partiallyspliced message, probes disposed upon EST arrays will often include multiple exons. The percentage of such exonspanning probes in an EST microarray can be calculated, on average, based upon the predicted number of exons/gene for the given species and the average length of the immobilized probes. For human genes, the near-complete sequence of human chromosome 22, Dunham et al., Nature 402(6761):489-95 (1999), predicts that human genes average 5.5 exons/gene. Even with probes of 200 - 500 bp, the vast majority of human EST microarray probes include more than one exon.

In contrast, by virtue of their origin from

algorithmically identified ORFs in genomic sequence, the probes in the genome-derived single exon microarrays of the present invention can consist of individual exons. Thus, in contrast to EST microarrays, at least about 50, 60, 70, 75, 80, 85, 95 or 99% of probes deposited in the genomederived microarray of the present invention consist of, or include, no more than one predicted ORF.

This provides the ability, not readily achieved using EST microarrays, to use the genome-derived single exon microarrays of the present invention to measure tissue-specific expression of individual exons, which in turn allows differential splicing events to be detected and characterized, and in particular, allows the correlation of differential splicing to tissue-specific expression patterns.

Furthermore, the exons that are represented in EST microarrays are often biased toward the 3' or 5' end of their respective genes, since sequencing strategies used for EST identification are so biased. In contrast, no such 3' or 5' bias necessarily inheres in the selection of exons for disposition on the genome-derived single exon microarrays of the present invention.

Conversely, the probes provided on the genomederived single exon microarrays of the present invention
typically, but need not necessarily, include intronic and/or intergenic sequence that is absent from EST microarrays, which are derived from mature mRNA.
Typically, at least about 50, 60, 70, 80 or 90% of the exon-including probes on the genome-derived single exon microarrays of the present invention include sequence drawn from noncoding regions. As discussed above, the additional presence of noncoding region does not significantly interfere with measurement of gene expression, and provides the additional opportunity to assay prespliced RNA, and thus measure such phenomena such as nuclear export control.

The genome-derived single exon microarrays of the present invention are also quite different from in situ synthesis microarrays, where probe size is severely constrained by inadequacies in the photolithographic synthesis process.

Typically, probes arrayed on in situ synthesis microarrays are limited to a maximum of about 25 bp. As a well known consequence, hybridization to such chips must be performed at low stringency. In order, therefore, to achieve unambiguous sequence-specific hybridization results, the in situ synthesis microarray requires substantial redundancy, with concomitant programmed arraying for each probe of probe analogues with altered (i.e., mismatched) sequence.

In contrast, the longer probe length of the genome-derived single exon microarrays of the present invention allows much higher stringency hybridization and wash. Typically, therefore, exon-including probes on the genome-derived single exon microarrays of the present invention average at least about 100, 200, 300, 400 or 500 bp in length. By obviating the need for substantial probe redundancy, this approach permits a higher density of probes for discrete exons or genes to be arrayed on the microarrays of the present invention than can be achieved for in situ synthesis microarrays.

A further distinction is that the probes in in situ synthesis microarrays typically are covalently linked to the substrate surface. In contrast, the probes disposed on the genome-derived microarray of the present invention typically are, but need not necessarily be, bound noncovalently to the substrate.

Furthermore, the short probe size on in situ microarrays causes large percentage differences in the melting temperature of probes hybridized to their complementary target sequence, and thus causes large

percentage differences in the theoretically optimum stringency across the array as a whole.

In contrast, the larger probe size in the microarrays of the present invention create lower percentage differences in melting temperature across the range of arrayed probes.

A further significant advantage of the microarrays of the present invention over in situ synthesized arrays is that the quality of each individual probe can be confirmed before deposition. In contrast, the quality of probes cannot be assessed on a probe-by-probe basis for the in situ synthesized microarrays presently being used.

The genome-derived single exon microarrays of the present invention are also distinguished over, and present substantial benefits over, the genome-derived microarrays from lower eukaryotes such as yeast. Lashkari et al., Proc. Natl. Acad. Sci. USA 94:13057-13062 (1997).

Only about 220 - 250 of the 6100 or so nuclear

genes in Saccharomyces cerevisiae - that is, only about 4
- 5% - have standard, spliceosomal, introns, Lopez et al.,
Nucl. Acids Res. 28:85-86 (2000); Spingola et al., RNA
5(2):221-34 (1999). Furthermore, the entire yeast genome
has already been sequenced. These two facts permit the
ready amplification and disposition of single-ORF amplicons
on such microarray without the requirement for antecedent
use of gene prediction and/or comparative sequence
analyses.

Thus, a significant aspect of the present

invention is the ability to identify and to confirm
expression of predicted coding regions in genomic sequence
drawn from eukaryotic organisms that have a higher
percentage of genes having introns than do yeast such as

Saccharomyces cerevisiae, particularly in genomic sequence
drawn from eukaryotes in which at least about 10, 20 or 50%

of protein-encoding genes have introns. In preferred embodiments, the methods and apparatus of the present invention are used to identify and confirm expression of novel genes from genomic sequence of eukaryotes in which the average number of introns per gene is at least about one, two or three or more.

. After the physical substrate is prepared, experimental verification of predicted function is performed.

In a preferred embodiment of the present invention, where the function sought to be identified in genomic sequence is protein coding, experimental verification is performed by measuring expression of the putative ORFs, typically through nucleic acid hybridization experiments, and in particularly preferred embodiments, through hybridization to genome-derived single exon microarrays prepared as above- described.

Expression is conveniently measured and expressed for each probe in the microarray as a ratio of the

20 expression measured concurrently in a plurality of mRNA sources, according to techniques well known in the microarray art, Reviewed in Schena et al., and as further described in Example 2, below. The mRNA source for the reference against which specific expression is measured can

25 be drawn from a homogeneous mRNA source, such as a single cultured cell-type, or alternatively can be heterogeneous, as from a pool of mRNA derived from multiple tissues and/or cell types, as further described in Example 2, infra.

mRNA can be prepared by standard techniques, see

30 Ausubel et al. and Maniatis et al., or purchased
commercially. The mRNA is then typically reversetranscribed in the presence of labeled nucleotides: the
index source (that in which expression is desired to be
measured) is reverse transcribed in the presence of

35 nucleotides labeled with a first label, typically a

fluorophore (fluorochrome; fluor; fluorescent dye); the reference source is reverse transcribed in the presence of a second label, typically a fluorophore, typically fluorometrically-distinguishable from the first label. As 5 further described in Example 2, infra, Cy3 and Cy5 dyes prove particularly useful in these methods. After partial purification of the index and reference targets, hybridization to the probe array is conducted according to standard techniques, typically under a coverslip.

After wash, microarrays are conveniently scanned using a commercial microarray scanning device, such as a Gen3 Scanner (Molecular Dynamics, Sunnyvale, CA). Data on expression is then passed, with or without interim storage, to process 500, where the results for each probe are 15 related to the original sequence.

10

Often, hybridization of target material to the genome-derived single exon microarray will identify certain of the probes thereon as of particular interest. Thus, it is often desirable that the user be able readily to obtain 20 sufficient quantities of an individual probe, either for subsequent arrayed deposition upon an additional support substrate, often as part of a microarray having a plurality of probes so identified, or alternatively or additionally as a solitary solid-phase or solution-phase probe, for 25 further use.

Thus, in another aspect, the present invention provides compositions and kits for the ready production of nucleic acids identical in sequence to, or substantially identical in sequence to, probes on the genome-derived 30 single exon microarrays of the present invention.

In this aspect, a small quantity of each probe is disposed, typically without attachment to substrate, in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate 35 can be used, greater efficiency is obtained using higher

density arrays, such as are provided by microtiter plates having 384, 864, 1536, 3456, 6144, or 9600 wells, and although microtiter plates having physical depressions (wells) are conveniently used, any device that permits addressable withdrawal of reagent from fluidly-noncommunicating areas can be used.

In this aspect of the invention, therefore, a fluidly noncommunicating addressable ordered set of individual probes, corresponding to those on a genome10 derived single exon microarray, is provided, with each probe in sufficient quantity to permit amplification, such as by PCR. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3'
15 primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this preferred embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes from the amplifiable ordered set.

Each discrete amplifiable probe can also be packaged with amplification primers, solutes, buffers, etc., and can be provided in dry (e.g., lyophilized) form or wet, in the latter case typically with addition of agents that retard evaporation.

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In another aspect of the present invention, a genome-derived single-exon microarray is packaged together with such an ordered set of amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In some embodiments, the microarray and/or ordered probe set are further packaged with recordable media that provide probe identification and addressing information, and that can additionally contain annotation

information, such as gene expression data. Such recordable media can be packaged with the microarray, with the ordered probe set, or with both.

If the microarray is constructed on a substrate that incorporates recordable media, such as is described in international patent application no. WO 98/12559, then separate packaging of the genome-derived single exon microarray and the bioinformatic information is not required.

The amount of amplifiable probe material should be sufficient to permit at least one amplification sufficient for subsequent hybridization assay.

Although the use of high density genome-derived microarrays on solid planar substrates is presently a preferred approach for the physical confirmation and characterization of the expression of sequences predicted to encode protein, other types of microarrays (as herein defined) can also be used.

Furthermore, as earlier mentioned, experimental verification of the function predicted from genomic sequence in process 200 can be bioinformatic, rather than, or additional to, physical verification.

For example, where the function desired to be identified is protein coding, the predicted ORFs can be compared bioinformatically to sequences known or suspected of being expressed.

Thus, the sequences output from process 300 (or process 200), can be used to query expression databases, such as EST databases, SNP ("single nucleotide") polymorphism") databases, known cDNA and mRNA sequences, SAGE ("serial analysis of gene expression") databases, and more generalized sequence databases that allow query for expressed sequences. Such query can be done by any sequence query algorithm, such as BLAST ("basic local alignment search tool"). The results of such query —

including information on identical sequences and
information on nonidentical sequences that have diffuse or
focal regions of sequence homology to the query sequence can then be passed directly to process 500, or used to
inform analyses subsequently undertaken in process 200,
process 300, or process 400.

Experimental data, whether obtained by physical or bioinformatic assay in process 400, is passed to process 500 where it is usefully related to the sequence data itself, a process colloquially termed "annotation". Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the record itself, by linking records in a hierarchical or relational database, by linking to external databases, or by a combination thereof. Such database techniques are well within the skill in the art.

The annotated sequence data can be stored locally, uploaded to genomic sequence database 100, and/or 20 displayed 800.

The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Coupled with the escalating pace at which sequence now accumulates, the rapid pace of sequence annotation produces a need for methods of displaying the information in meaningful ways.

FIG. 3 shows visual display 80 presenting a single genomic sequence annotated according to the present invention. Because of its nominal resemblance to artistic works of Piet Mondrian, visual display 80 is alternatively described herein as a "Mondrian".

Each of the visual elements of display 80 is aligned with respect to the genomic sequence being annotated (hereinafter, the "annotated sequence"). Given the number of nucleotides typically represented in an

annotated sequence, representation of individual nucleotides would rarely be readable in hard copy output of display 80. Typically, therefore, the annotated sequence is schematized as rectangle 89, extending from the left 5 border of display 80 to its right border. By convention herein, the left border of rectangle 89 represents the first nucleotide of the sequence and the right border of rectangle 89 represents the last nucleotide of the sequence.

10

As further discussed below, however, the Mondrian visual display of annotated sequence can serve as a convenient graphical user interface for computerized representation, analysis, and query of information stored electronically. For such use, the individual nucleotides 15 can conveniently be linked to the X axis coordinate of rectangle 89. This permits the annotated sequence at any point within rectangle 89 readily to be viewed, either automatically - for example, by time-delayed appearance of a small overlaid window upon movement of a cursor or other 20 pointer over rectangle 89 - or through user intervention, as by clicking a mouse or other pointing device at a point in rectangle 89.

Visual display 80 is generated after user specification of the genomic sequence to be displayed. 25 Such specification can consist of or include an accession number for a single clone (e.g., a single BAC accessioned into GenBank), wherein the starting and stopping nucleotides are thus absolutely identified, or alternatively can consist of or include an anchor or 30 fulcrum point about which a chosen range of sequence is anchored, thus providing relative endpoints for the sequence to be displayed. For example, the user can anchor such a range about a given chromosomal map location, gene name, or even a sequence returned by query for similarity 35 or identity to an input query sequence. When visual

display 80 is used as a graphical user interface to computerized data, additional control over the first and last displayed nucleotide will typically be dynamically selectable, as by use of standard zooming and/or selection tools.

Field 81 of visual display 80 is used to present the output from process 200, that is, to present the bioinformatic prediction of those sequences having the desired function within the genomic sequence. Functional sequences are typically indicated by at least one rectangle 83 (83a, 83b, 83c), the left and right borders of which respectively indicate, by their X-axis coordinates, the starting and ending nucleotides of the region predicted to have function.

Where a single bioinformatic method or approach identifies a plurality of regions having the desired function, a plurality of rectangles 83 is disposed horizontally in field 81. Where multiple methods and/or approaches are used to identify function, each such method and/or approach can be represented by its own series of horizontally disposed rectangles 83, each such horizontally disposed series of rectangles offset vertically from those representing the results of the other methods and approaches.

25 Thus, rectangles 83a in FIG. 3 represent the functional predictions of a first method of a first approach for predicting function, rectangles 83b represent the functional predictions of a second method and/or second approach for predicting that function, and rectangles 83c represent the predictions of a third method and/or approach.

Where the function desired to be identified is protein coding, field 81 is used to present the bioinformatic prediction of sequences encoding protein.

For example, rectangles 83a can represent the results from

GRAIL or GRAIL II, rectangles 83b can represent the results from GENEFINDER, and rectangles 83c can represent the results from DICTION.

Optionally, and preferably, rectangles 83

5 collectively representing predictions of a single method and/or approach are identically colored and/or textured, and are distinguishable from the color and/or texture used for a different method and/or approach.

Alternatively, or in addition, the color, hue,

density, or texture of rectangles 83 can be used further to
report a measure of the bioinformatic reliability of the
prediction. For example, many gene prediction programs
will report a measure of the reliability of prediction.
Thus, increasing degrees of such reliability can be

indicated, e.g., by increasing density of shading. Where
display 80 is used as a graphical user interface, such
measures of reliability, and indeed all other results
output by the program, can additionally or alternatively be
made accessible through linkage from individual rectangles

83, as by time-delayed window ("tool tip" window), or by
pointer (e.g., mouse)-activated link.

As earlier described, increased predictive reliability can be achieved by requiring consensus among methods and/or approaches to determining function. Thus, field 81 can include a horizontal series of rectangles 83 that indicate one or more degrees of consensus in predictions of function.

Although FIG. 3 shows three series of horizontally disposed rectangles in field 81, display 80 can include as few as one such series of rectangles and as many as can discriminably be displayed, depending upon the number of methods and/or approaches used to predict a given function.

Furthermore, field 81 can be used to show predictions of a plurality of different functions.

However, the increased visual complexity occasioned by such display makes more useful the ability of the user to select a single function for display. When display 80 is used as a graphical user interface for computer query and analysis, such function can usefully be indicated and userselectable, as by a series of graphical buttons or tabs (not shown in FIG. 3).

Rectangle 89 is shown in FIG. 3 as including interposed rectangle 84. Rectangle 84 represents the 10 portion of annotated sequence for which predicted functional information has been assayed physically, with the starting and ending nucleotides of the assayed material indicated by the X axis coordinates of the left and right borders of rectangle 84. Rectangle 85, with optional 15 inclusive circles 86 (86a, 86b, and 86c) displays the results of such physical assay.

Although a single rectangle 84 is shown in FIG. 3, physical assay is not limited to just one region of annotated genomic sequence. It is expected that an increasing percentage of regions predicted to have function by process 200 will be assayed physically, and that display 80 will accordingly, for any given genomic sequence, have an increasing number of rectangles 84 and 85, representing an increased density of sequence annotation.

Where the function desired to be identified is protein coding, rectangle 84 identifies the sequence of the probe used to measure expression. In embodiments of the present invention where expression is measured using genome-derived single exon microarrays, rectangle 84 identifies the sequence included within the probe immobilized on the support surface of the microarray. As noted supra, such probe will often include a small amount of additional, synthetic, material incorporated during amplification and designed to permit reamplification of the probe, which sequence is typically not shown in display 80.

Rectangle 87 is used to present the results of bioinformatic assay of the genomic sequence. For example, where the function desired to be identified is protein coding, process 400 can include bioinformatic query of 5 expression databases with the sequences predicted in process 200 to encode exons. And as earlier discussed, because bioinformatic assay presents fewer constraints than does physical assay, often the entire output of process 200 can be used for such assay, without further subsetting 10 thereof by process 300. Therefore, rectangle 87 typically need not have separate indicators therein of regions submitted for bioinformatic assay; that is, rectangle 87 typically need not have regions therein analogous to rectangles 84 within rectangle 89.

Rectangle 87 as shown in FIG. 3 includes smaller rectangles 880 and 88. Rectangles 880 indicate regions that returned a positive result in the bioinformatic assay, with rectangles 88 representing regions that did not return such positive results. Where the function desired to be 20 predicted and displayed is protein coding, rectangles 880 indicate regions of the predicted exons that identify sequence with significant similarity in expression databases, such as EST, SNP, SAGE databases, with rectangles 88 indicating genes novel over those identified 25 in existing expression data bases.

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Rectangles 880 can further indicate, through color, shading, texture, or the like, additional information obtained from bioinformatic assay.

For example, where the function assayed and 30 displayed is protein coding, the degree of shading of rectangles 880 can be used to represent the degree of sequence similarity found upon query of expression databases. The number of levels of discrimination can be as few as two (identity, and similarity, where similarity 35 has a user-selectable lower threshold). Alternatively, as

many different levels of discrimination can be indicated as can visually be discriminated.

Where display 80 is used as a graphical user interface, rectangles 880 can additionally provide links directly to the sequences identified by the query of expression databases, and/or statistical summaries thereof. As with each of the precedingly-discussed uses of display 80 as a graphical user interface, it should be understood that the information accessed via display 80 need not be resident on the computer presenting such display, which often will be serving as a client, with the linked information resident on one or more remotely located servers.

Rectangle 85 displays the results of physical assay of the sequence delimited by its left and right borders.

Rectangle 85 can consist of a single rectangle, thus indicating a single assay, or alternatively, and increasingly typically, will consist of a series of rectangles (85a, 85b, 85c) indicating separate physical assays of the same sequence.

Where the function assayed is gene expression, and where gene expression is assayed as herein described using simultaneous two-color fluorescent detection of hybridization to genome-derived single exon microarrays, individual rectangles 85 can be colored to indicate the degree of expression relative to control. Conveniently, shades of green can be used to depict expression in the sample over control values, and shades of red used to depict expression less than control, corresponding to the spectra of the Cy3 and Cy5 dyes conventionally used for respective labeling thereof. Additional functional information can be provided in the form of circles 86 (86a, 86b, 86c), where the diameter of the circle can be used to indicate expression intensity. As discussed infra, such

relative expression (expression ratios) and absolute expression (signal intensity) can be expressed using normalized values.

Where display 80 is used as a graphical user

5 interface, rectangle 85 can be used as a link to further
information about the assay. For example, where the assay
is one for gene expression, each rectangle 85 can be used
to link to information about the source of the hybridized
mRNA, the identity of the control, raw or processed data

10 from the microarray scan, or the like.

FIG. 4 is rendition of display 80 representing gene prediction and gene expression for a hypothetical BAC, showing conventions used in the Examples presented infra.

BAC sequence ("Chip seq.") 89 is presented, with the physically assayed region thereof (corresponding to rectangle 84 in FIG. 3) shown in white. Algorithmic gene predictions are shown in field 81, with predictions by GRAIL shown, predictions by GENEFINDER, and predictions by DICTION shown. Within rectangle 87, regions of sequence that, when used to query expression databases, return identical or similar sequences ("EST hit") are shown as white rectangles (corresponding to rectangles 880 in FIG. 3), gray indicates low homology, and black indicates unknowns (where black and gray would correspond to rectangles 88 in FIG. 3).

Although FIGS. 3 and 4 show a single stretch of sequence, uninterrupted from left to right, longer sequences are usefully represented by vertical stacking of such individual Mondrians, as shown in FIGS. 9 and 10.

Single Exon Probes Useful For Measuring Gene Expression

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The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Where the function to be identified is

protein coding, the methods and apparatus of the present invention rapidly identify and confirm the expression of portions of genomic sequence that function to encode protein. As a direct result, the methods and apparatus of the present invention rapidly yield large numbers of single-exon nucleic acid probes, the majority from previously unknown genes, each of which is useful for measuring and/or surveying expression of a specific gene in one or more tissues or cell types.

It is, therefore, another aspect of the present invention to provide genome-derived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray.

Using the methods and genome-derived single-exon

15 microarrays of the present invention, we have for example readily identified a large number of unique ORFs from human genomic sequence. Using single exon probes that encompass these ORFs, we have demonstrated, through microarray hybridization analysis, the expression of 12,821 of these

20 ORFs in brain.

As would immediately be appreciated by one of skill in the art, each single exon probe having demonstrable expression in brain is currently available for use in measuring the level of its ORF's expression in brain.

Diseases of the brain and nervous system are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the brain and nervous system, for the most part these disorders are believed to have polygenic etiologies.

For example, over the past few decades

35 Alzheimer's disease (AD), once considered a rare disorder,

has become recognized as a major public health problem; over 4,000,000 people in the United States are now estimated to suffer with various stages of this progressive, degenerative brain disorder.

Although there is no agreement on the exact incidence or prevalence of Alzheimer's disease, in part due to varying diagnostic criteria and difficulties of differential diagnosis among dementias, the studies are consistent in pointing to an exponential rise in prevalence of this disease with age. After age 65, the percentage of affected people approximately doubles with every decade of life, regardless of definition. Among people age 85 or older, studies suggest that 25 to 35 percent have dementia, including Alzheimer's disease; one study reports that 47.2 percent of people over age 85 have Alzheimer's disease, exclusive of other dementias.

Alzheimer's disease progressively destroys memory, reason, judgment, language, and, eventually, the ability to carry out even the simplest of tasks. Anatomic changes associated with Alzheimer's disease begin in the entorhinal cortex, proceed to the hippocampus, and then gradually spread to other regions, particularly the cerebral cortex. Chief among such anatomic changes are the presence of characteristic extracellular plaques and internal neurofibrillary tangles.

Alzheimer's disease has been suspected to have a multifactorial genetic etiological component for almost half a century. Sjogren et al., Acta Psychiat. Neurol. Scand. 82(suppl.): 1-152 (1952).

At least four genes have been identified to date that contribute to development of Alzheimer's disease: AD1 is caused by mutations in the amyloid precursor gene (APP); AD2 is associated with the APOE4 allele on chromosome 19; AD3 is caused by mutation in a chromosome 14 gene encoding a 7-transmembrane domain protein, presentlin-1 (PSEN1), and

AD4 is caused by mutation in a gene on chromosome 1 that encodes a similar 7-transmembrane domain protein, presentlin-2 (PSEN2).

There is strong evidence, however, for additional, as yet uncharacterized, AD loci on other chromosomes.

For example, Daw et al., Am. J. Hum. Genet. 66: 196-204 (2000), estimated the number of additional quantitative trait loci (QTLs) and their contribution to the variance in age at onset of AD, and reported that 4 loci make a contribution to the variance in age at onset of late-onset AD similar to or greater in magnitude than that made by apoE, with one locus making a contribution several times greater than that of apoE. These results suggest that several genes not yet localized may play a larger role than does apoE in late-onset AD.

In accord, three groups recently announced the possible existence of an AD susceptibility gene on chromosome 10. Bertram et al., Science 290(5500):2302-2303 (2000); Ertekin-Taner et al., Science 290(5500):2303-2304 (2000); and Myers et al., Science 290(5500):2304-23055 (2000).

As another example, multiple sclerosis (MS)

25 affects about 350,000 Americans, with approximately 200 new cases diagnosed each week, with an estimated annual monetary cost in the U.S. alone of \$2.5 billion.

Clinically, MS is an unpredictable disorder, with symptoms, presentation and course falling broadly into one of several clinical patterns. In relapsing-remitting (RR) MS, the disease first manifests as a series of attacks followed by complete or partial remissions, with symptoms returning later after a period of stability. In primary-progressive (PP) MS, there is a gradual clinical decline with no distinct remissions, although there may be

temporary plateaus or minor relief from symptoms.

Secondary-progressive (SP) MS begins with a relapsingremitting course followed by a later primary-progressive
course. Rarely, patients may have a progressive-relapsing

(PR) course in which the disease takes a progressive
path punctuated by acute attacks. PP, SP, and PR MS are
sometimes lumped together and called chronic progressive
MS. The waxing and waning course characteristic of RR, SP
and PR MS makes differential diagnosis difficult.

Anatomically, MS attacks are associated with focal inflammation in areas of the white matter of the central nervous system (CNS), accompanied or followed by demyelination in these areas, termed plaques. Destruction of the myelin sheath slows or blocks neurological transmission, leading to diminished or lost function. Clinical manifestations depend upon the location of the plaques and severity of demyelination, and range from fatigue, the most common symptom of MS, to visual impairment, due to inflammation of the optic nerve, termed optic neuritis, to numbness and paresthesias, to focal muscular weakness, ataxia, and bladder incontinence.

Increasing evidence suggests that genotype contributes to susceptibility to MS.

As early as 1965, McAlpine, in Multiple

25 Sclerosis: A Reappraisal (McAlpine, ed.), Williams and
Wilkins Co. pp. 61-74 (1965), concluded that the risk to a
first-degree relative of a patient with multiple sclerosis
is at least 15 times that for a member of the general
population, but could discern no definite genetic pattern

30 of inheritance.

Subsequently, many studies associated MS with HLA (MHC) haplotype. Haines et al., Hum. Molec. Genet. 7:1229-1234 (1998), studying a data set of 98 multiplex MS families, confirmed earlier reports that genetic linkage to the MHC can be explained by association with the HLA-DR2

allele, but suggested that MHC association explains only between 17% and 62% of the genetic etiology of MS.

From a review of genomic screens, Dyment et al., Hum. Molec. Genet. 6: 1693-1698 (1997), concluded that a 5 number of genes with interacting effects are likely and that no single region has a major influence on familial risk. Chataway et al., Brain 121: 1869-1887 (1998), reporting a follow-up on U.K. studies using a systematic genome screen to determine the genetic basis of MS, stated 10 that a gene of major effect had been excluded from 95% of the genome and one with a moderate role from 65%, results thus suggesting that multiple sclerosis depends on independent or epistatic effects of several genes, each with small individual effects, rather than a very few genes 15 of major biologic importance.

As a yet further example, schizophrenia has long been recognized to have complex, likely polygenic, genetic contributions.

Schizophrenia is a common psychiatric disorder, 20 occurring in 1 to 1.5 percent of the population worldwide, and is characterized by variable constellations of symptoms drawn from a universe of behavioral abnormalities. Although there are accepted alternative diagnostic criteria, primary criteria for diagnosis require two or 25 more of the following, each present for a significant portion of time during a 1-month period (or less if successfully treated): (1) delusions; (2) hallucinations ; (3) disorganized speech (e.g., frequent derailment or incoherence); (4) grossly disorganized or catatonic 30 behavior; (5) negative symptoms, i.e., affective flattening, alogia, or avolition. (Diagnostic and Statistic Manual of Mental Disorders DSM-IV-TR, American Psychiatric Association (2000)). Only one such symptom is required if delusions are bizarre or hallucinations

35 consist of a voice keeping up a running commentary on the

person's behavior or thoughts, or consist of two or more voices conversing with each other.

Three-quarters of persons with schizophrenia develop the disease between 16 and 25 years of age: onset is uncommon after age 30, rare after age 40. In the 16 to 25 year old age group, schizophrenia affects more men than women; in the 25-30 year old group, the incidence is higher in women than in men. Studies have shown that some persons with schizophrenia recover completely, and many others improve to the point where they can live independently, often with the maintenance of drug therapy. However, approximately 15 percent of people with schizophrenia respond only moderately to medication and require extensive support throughout their lives, while another 15 percent simply do not respond to existing treatment.

Schizophrenia has long been known to have a significant genetic component. Studies have consistently demonstrated that the risk to relatives of a proband with schizophrenia is higher than the risk to relatives of 20 controls. Moldin, in Genetics and Mental Disorders: Report of the NIMH Genetics Workgroup (NIH publication 98-4268, (1998), reviewed family and twin studies published between 1920 and 1987 and found the recurrence risk ratios to be 48 for monozygotic twins, 11 for first-degree relatives, 4.25 25 for second-degree relatives, and 2 for third-degree relatives. He also found that concordance rates for monozygotic twins averaged 46%, even when reared in different families, whereas the concordance rates for dizygotic twins averaged only 14%. The prevalence of 30 schizophrenia is known to be higher in biologic than in adoptive relatives of schizophrenic adoptees.

The mode of inheritance is unclear, however.

Susceptibility has been mapped to many loci, including chromosomes 1q21-q22, 5, 6p23, 8p22-p21, 11q, 13q14-q21, 13q32, 15q15, 15q14, 18p, and 22q11. Chromosome

19 has also been implicated in schizophrenia, at 2 different sites, as have sites on the X chromosome. Wei et al., Nature Genet. 25:376-377 (2000) report more specifically that the NOTCH4 locus is associated with susceptibility to schizophrenia.

In general, however, it is believed that development of schizophrenia involves multiple loci.

For example, Williams et al., Hum. Molec. Genet. 8:1729-1739 (1999) undertook a systematic search for linkage in 196 affected sib pairs (ASPs) with schizophrenia. Using 229 microsatellite markers at an average intermarker distance of 17.26 cM, followed in a second stage by a further 54 markers allowing the regions identified in stage 1 to be typed at an average spacing of 5.15 cM, Williams et al. considered results on chromosomes 4p, 18q, and Xcen as suggestive; however, given the scores, Williams et al. interpreted their results as suggesting that common genes of major effect (susceptibility ratio more than 3) are unlikely to exist for schizophrenia.

Similarly, Shaw et al., Am. J. Med. Genet. 81(5):364-76 (1998), in a genome-wide search for schizophrenia susceptibility genes, found that twelve chromosomes (1, 2, 4, 5, 8, 10, 11, 12, 13, 14, 16, and 22) had at least one region with a nominal P value <0.05, that two of these chromosomes had a nominal P value <0.01 (chromosomes 13 and 16), and that five chromosomes (1, 2, 4, 11, and 13) had at least one marker with a lod score >2.0, suggesting the existence of multiple loci that contribute to schizophrenia susceptibility.

As yet another example, multiple genes are thought to predispose to epilepsy.

Epilepsy is characterized by recurrent, paroxysmal disorders of cerebral function (seizures); that is, by sudden, brief attacks of altered consciousness, motor activity, sensory phenomena, or inappropriate

behavior. The risk of developing epilepsy is 1% in the period from birth to age 20, and 3% at age 75.

Epilepsy is caused by excessive discharge of cerebral neurons. Clinical manifestations depend on the type and location of discharge. In partial seizures, for example, the excess neuronal discharge is contained within one region of the cerebral cortex. Simple partial seizures consist of motor, sensory, or psychomotor phenomena without loss of consciousness; the specific phenomenon reflects the affected area of the brain. In generalized seizures, the discharge bilaterally and diffusely involves the entire cortex. Sometimes a focal lesion of one part of a hemisphere activates the entire cerebrum bilaterally so rapidly that it produces a generalized tonic-clonic seizure before a focal sign appears.

Epilepsy is a family of disorders. Those that are idiopathic are believed to have multiple genetic contributions. For example, idiopathic generalized epilepsy (IGE) is characterized by recurring generalized seizures in the absence of detectable brain lesions and/or metabolic abnormalities. Twin and family studies suggest that genetic factors play a key part in its etiology. Although a mutation in the CACNB4 gene can cause the disorder, linkage to 8q24, Zara et al., Hum. Molec. Genet. 4: 1201-1207(1995), 3q26 and 14q23, Sander et al., Hum. Molec. Genet. 9:1465-1472 (2000), and 2q36 has been also demonstrated, with a multilocus model appearing to fit best the observed familial patterns.

Polygenic contributions to the etiology of various neurologic cancers have similarly been described.

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For example, gliomas account for 45% of intracranial tumors, and multiple loci have been implicated in its development, with losses of chromosome 17p, increase in copy number of chromosome 7, structural abnormalities of

chromosomes 9p and 19q, and genes on chromosome 10 among the suspects.

Other significant diseases of brain and nervous tissue are also believed to have a genetic, typically 5 polygenic, etiologic component. These diseases include, for example, Parkinson's disease, dementia with Lewy bodies, frontotemporal dementia, corticobasal ganglionic degeneration, progressive supranuclear palsy, prion diseases (Creutzfeld-Jakob, Gerstmann-Strausller-Shenker, 10 familial fatal insomnia), Tourette's Syndrome, corticobasal degeneration, multiple system atrophy, striatonigral degeneration, Shy-Drager syndrome, olivopontocerebellar atrophy, spinocerebellar ataxia, Friedreich ataxia, ataxiatelangiectasia, amyotrophic lateral sclerosis, bulbospinal 15 atrophy (Kennedy's syndrome), spinal muscular atrophy, neuronal storage diseases (sphingolipid, mucopolysaccharide, mucolipid), leukodystrophy, Krabbe disease, metachromic leukodystrophy, adrenoleukodystrophy, Pelizaeus-Merzbacher disease, Canavan disease, 20 mitochondrial encephalomyopathy, Leigh disease, neurofibromatosis (Type 1 and Type II), tuberous sclerosis, paraneoplastic syndrome, subacute cerebellar degeneration,

subacute sensory neuropathy, opsoclonus/myoclonus, retinal degeneration, stiff-man syndrome and Von Hippel-Lindau 25 disease.

Many neurologic cancers other than gliomas have also been shown or suspected to have genetic bases or contributions. Among these cancers are astrocytoma, fibrillary astrocytoma, pilocytic astrocytoma,

30 pleomorphic xanthoastrocytoma, oligodendroglioma, ependymoma, gangliocytoma, ganglioglioma, medulloblastoma, primary brain germ cell tumor, pineocytoma, pineoblastoma, and meningioma.

Other disorders of brain and central nervous 35 system that likely have genetic components include the

various forms of neural deafness, catatonia, depression, bipolar (manic-depressive) disorder, Wilson's Disease, Pick disease, neuromyelitis optica (Devic disease), central pontine myelinolysis, Marchiafava-Bignami disease, 5 Guillain-Barre syndrome, sleep disorders (insomnia, myoclonus, narcolepsy, cataplexy, sleep apnea), amnesia, aphasias (including Broca's aphasia and Wernicke's aphasia), cortical blindness, visual agnosia, auditory agnosia, and Kluver-Bucy syndrome.

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The human genome-derived single exon nucleic acid probes and microarrays of the present invention are useful for predicting, diagnosing, grading, staging, monitoring and prognosing diseases of human brain, particularly those diseases with polygenic etiology. With each of the single 15 exon probes described herein shown to be expressed at detectable levels in human brain, and with about 2/3 of the probes identifying novel genes, the single exon microarrays of the present invention provide exceptionally high informational content for such studies.

For example, diagnosis (including differential diagnosis among clinically indistinguishable disorders), staging, and/or grading of a disease can be based upon the quantitative relatedness of a patient gene expression profile to one or more reference expression profiles known 25 to be characteristic of a given neurologic disease, or to specific grades or stages thereof.

In one embodiment, the patient gene expression profile is generated by hybridizing nucleic acids obtained directly or indirectly from transcripts expressed in the 30 patient's brain (or other CNS tissues, including cultured tissues) to the genome-derived single exon microarray of the present invention. Reference profiles are be obtained similarly by hybridizing nucleic acids from individuals with known disease. Methods for quantitatively relating 35 gene expression profiles, without regard to the function of

the protein encoded by the gene, are disclosed in WO 99/58720, incorporated herein by reference in its entirety.

In another approach, the genome-derived single exon probes and microarrays of the present invention can be used to interrogate genomic DNA, rather than pools of expressed message; this latter approach permits predisposition to and/or prognosis of neurologic disease to be assessed through the massively parallel determination of altered copy number, deletion, or mutation in the patient's genome of exons known to be expressed in human brain. The algorithms set forth in WO 99/58720 can be applied to such genomic profiles without regard to the function of the protein encoded by the interrogated gene.

The utility is specific to the probe; at

15 sufficiently high hybridization stringency, which

stringencies are well known in the art — see Ausubel et al.

and Maniatis et al. — each probe reports the level of

expression of message specifically containing that ORF.

It should be appreciated, however, that the
20 probes of the present invention, for which expression in
the brain has been demonstrated are useful for both
measurement in the brain and for survey of expression in
other tissues.

Significant among such advantages is the presence of probes for novel genes.

As mentioned above and further detailed in Examples 1 and 2, the methods described enable ORFs which are not present in existing expression databases to be identified. And the fewer the number of tissues in which the ORF can be shown to be expressed, the more likely the ORF will prove to be part of a novel gene: as further discussed in Example 2, ORFs whose expression was measurable in only a single of the tested tissues were represented in existing expression databases at a rate of only 11%, whereas 36% of ORFs whose expression was

measurable in 9 tissues were present in existing expression databases, and fully 45% of those ORFs expressed in all ten tested tissues were present in existing expressed sequence databases.

Either as tools for measuring gene expression or tools for surveying gene expression, the genome-derived single exon probes of the present invention have significant advantages over the cDNA or EST-based probes that are currently available for achieving these utilities.

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The genome-derived single exon probes of the present invention are useful in constructing genome-derived single exon microarrays; the genome-derived single exon microarrays, in turn, are useful devices for measuring and for surveying gene expression in the human.

Gene expression analysis using microarrays — conventionally using microarrays having probes derived from expressed message — is well-established as useful in the biological research arts (see Lockhart et al. Nature 405, 827-836).

20 Microarrays have been used to determine gene expression profiles in cells in response to drug treatment (see, for example, Kaminski et al., "Global Analysis of Gene Expression in Pulmonary Fibrosis Reveals Distinct Programs Regulating Lung Inflammation and Fibrosis," Proc.

25 Natl. Acad. Sci. USA 97(4):1778-83 (2000); Bartosiewicz et al., "Development of a Toxicological Gene Array and Quantitative Assessment of This Technology," Arch. Biochem. Biophys. 376(1):66-73 (2000)), viral infection (see for example, Geiss et al., "Large-scale Monitoring of Host Cell Gene Expression During HIV-1 Infection Using cDNA Microarrays," Virology 266(1):8-16 (2000)) and during cell processes such as differentiation, senescence and apoptosis (see, for example, Shelton et al., "Microarray Analysis of

35 Voehringer et al., "Gene Microarray Identification of Redox

Replicative Senescence, " Curr. Biol. 9(17):939-45 (1999);

and Mitochondrial Elements That Control Resistance or Sensitivity to Apoptosis," Proc. Natl. Acad. Sci. USA 97(6):2680-5 (2000)).

Microarrays have also been used to determine 5 abnormal gene expression in diseased tissues (see, for example, Alon et al., "Broad Patterns of Gene Expression Revealed by Clustering Analysis of Tumor and Normal Colon Tissues Probed by Oligonucleotide Arrays," Proc. Natl. Acad. Sci. USA 96(12):6745-50 (1999); Perou et al., 10 "Distinctive Gene Expression Patterns in Human Mammary Epithelial Cells and Breast Cancers, Proc. Natl. Acad. Sci. USA 96(16):9212-7 (1999); Wang et al., "Identification of Genes Differentially Over-expressed in Lung Squamous Cell Carcinoma Using Combination of cDNA Subtraction and 15 Microarray Analysis," Oncogene 19(12):1519-28 (2000); Whitney et al., "Analysis of Gene Expression in Multiple Sclerosis Lesions Using cDNA Microarrays," Ann. Neurol. 46(3):425-8 (1999)), in drug discovery screens (see, for example, Scherf et al., "A Gene Expression Database for the

20 Molecular Pharmacology of Cancer," Nat. Genet. 24(3):236-44 (2000)) and in diagnosis to determine appropriate treatment strategies (see, for example, Sgroi et al., "In vivo Gene Expression Profile Analysis of Human Breast Cancer Progression," Cancer Res. 59(22):5656-61 (1999)).

In microarray-based gene expression screens of pharmacological drug candidates upon cells, each probe provides specific useful data. In particular, it should be appreciated that even those probes that show no change in expression are as informative as those that do change, serving, in essence, as negative controls.

For example, where gene expression analysis is used to assess toxicity of chemical agents on cells, the failure of the agent to change a gene's expression level is evidence that the drug likely does not affect the pathway of which the gene's expressed protein is a part.

Analogously, where gene expression analysis is used to assess side effects of pharmacological agents - whether in lead compound discovery or in subsequent screening of lead compound derivatives - the inability of the agent to alter 5 a gene's expression level is evidence that the drug does not affect the pathway of which the gene's expressed protein is a part.

WO 99/58720 provides methods for quantifying the relatedness of a first and second gene expression profile 10 and for ordering the relatedness of a plurality of gene expression profiles. The methods so described permit useful information to be extracted from a greater percentage of the individual gene expression measurements from a microarray than methods previously used in the art.

Other uses of microarrays are described in Gerhold et al., Trends Biochem. Sci. 24(5):168-173 (1999) and Zweiger, Trends Biotechnol. 17(11):429-436 (1999); Schena et al.

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The invention particularly provides genome-20 derived single-exon probes known to be expressed in brain.

The individual single exon probes can be provided in the form of substantially isolated and purified nucleic acid, typically, but not necessarily, in a quantity sufficient to perform a hybridization reaction.

Such nucleic acid can be in any form directly hybridizable to the message that contains the probe's ORF, such as double stranded DNA, single-stranded DNA complementary to the message, single-stranded RNA complementary to the message, or chimeric DNA/RNA molecules 30 so hybridizable. The nucleic acid can alternatively or additionally include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained. For example, probes can include phosphorothioates, methylphosphonates,

35 morpholino analogs, and peptide nucleic acids (PNA), as are

described, for example, in U.S. Patent Nos. 5,142,047; 5,235,033; 5,166,315; 5,217,866; 5,184,444; 5,861,250.

Usefully, however, such probes are provided in a form and quantity suitable for amplification, where the samplified product is thereafter to be used in the hybridization reactions that probe gene expression. Typically, such probes are provided in a form and quantity suitable for amplification by PCR or by other well known amplification technique. One such technique additional to PCR is rolling circle amplification, as is described, inter alia, in U.S. Patent Nos. 5,854,033 and 5,714,320 and international patent publications WO 97/19193 and WO 00/15779. As is well understood, where the probes are to be provided in a form suitable for amplification, the range of nucleic acid analogues and/or internucleotide linkages will be constrained by the requirements and nature of the amplification enzyme.

Where the probe is to be provided in form suitable for amplification, the quantity need not be sufficient for direct hybridization for gene expression analysis, and need be sufficient only to function as an amplification template, typically at least about 1, 10 or 100 pg or more.

Each discrete amplifiable probe can also be

25 packaged with amplification primers, either in a single
composition that comprises probe template and primers, or
in a kit that comprises such primers separately packaged
therefrom. As earlier mentioned, the ORF-specific
5' primers used for genomic amplification can have a first

30 common sequence added thereto, and the ORF-specific 3'
primers used for genomic amplification can have a second,
different, common sequence added thereto, thus permitting,
in this embodiment, the use of a single set of 5' and 3'
primers to amplify any one of the probes. The probe

35 composition and/or kit can also include buffers, enzyme,

غر):

etc., required to effect amplification.

As mentioned earlier, when intended for use on a genome-derived single exon microarray of the present invention, the genome-derived single exon probes of the present invention will typically average at least about 100, 200, 300, 400 or 500 bp in length, including (and typically, but not necessarily centered about) the ORF. Furthermore, when intended for use on a genome-derived single exon microarray of the present invention, the genome-derived single exon probes of the present invention will typically not contain a detectable label.

When intended for use in solution phase
hybridization, however — that is, for use in a
hybridization reaction in which the probe is not first
bound to a support substrate (although the target may
indeed be so bound) — length constraints that are imposed
in microarray-based hybridization approaches will be
relaxed, and such probes will typically be labeled.

In such case, the only functional constraint that

dictates the minimum size of such probe is that each such

probe must be capable of specifically identifying in a

hybridization reaction the exon from which it is drawn. In

theory, a probe of as little as 17 nucleotides is capable

of uniquely identifying its cognate sequence in the human

genome. For hybridization to expressed message — a subset

of target sequence that is much reduced in complexity as

compared to genomic sequence — even fewer nucleotides are

required for specificity.

Therefore, the probes of the present invention

on include as few as 20, 25 or 50 bp or ORF, or more. In particular embodiments, the ORF sequences are given in SEQ ID NOS. 12,822 - 25,434, respectively, for probe SEQ ID NOS. 1 - 12,821. The minimum amount of ORF required to be included in the probe of the present invention in order to provide specific signal in either solution phase or

microarray-based hybridizations can readily be determined for each of ORF SEQ ID NOS. 12,822 - 25,434 individually by routine experimentation using standard high stringency conditions.

5 Such high stringency conditions are described, inter alia, in Ausubel et al. and Maniatis et al. For microarray-based hybridization, standard high stringency conditions can usefully be 50% formamide, 5X SSC, 0.2 $\mu g/\mu l$ poly(dA), 0.2 $\mu g/\mu l$ human cot1 DNA, and 0.5 % SDS, in a 10 humid oven at 42°C overnight, followed by successive washes of the microarray in 1X SSC, 0.2% SDS at 55°C for 5 $\,$ minutes, and then 0.1% SSC, 0.2% SDS, at 55°C for 20 $\,$ minutes. For solution phase hybridization, standard high stringency conditions can usefully be aqueous hybridization 15 at 65°C in 6X SSC. Lower stringency conditions, suitable for cross-hybridization to mRNA encoding structurally- and functionally-related proteins, can usefully be the same as the high stringency conditions but with reduction in temperature for hybridization and washing to room 20 temperature (approximately 25°C).

When intended for use in solution phase hybridization, the maximum size of the single exon probes of the present invention is dictated by the proximity of other expressed exons in genomic DNA: although each single exon probe can include intergenic and/or intronic material contiguous to the ORF in the human genome, each probe of the present invention will include portions of only one expressed exon.

Thus, each single exon probe will include no more than about 25 kb of contiguous genomic sequence, more typically no more than about 20 kb of contiguous genomic sequence, more usually no more than about 15 kb, even more usually no more than about 10 kb. Usually, probes that are maximally about 5 kb will be used, more typically no more than about 3 kb.

It will be appreciated that the Sequence Listing appended hereto presents, by convention, only that strand of the probe and ORF sequence that can be directly translated reading from 5' to 3' end. As would be well understood by one of skill in the art, single stranded probes must be complementary in sequence to the ORF as present in an mRNA; it is well within the skill in the art to determine such complementary sequence. It will further be understood that double stranded probes can be used in both solution-phase hybridization and microarray-based hybridization if suitably denatured.

Thus, it is an aspect of the present invention to provide single-stranded nucleic acid probes that have sequence complementary to those described herein above and below, and double-stranded probes one strand of which has sequence complementary to the probes described herein.

The probes can, but need not, contain intergenic and/or intronic material that flanks the ORF, on one or both sides, in the same linear relationship to the ORF that the intergenic and/or intronic material bears to the ORF in genomic DNA. The probes do not, however, contain nucleic acid derived from more than one expressed ORF.

And when intended for use in solution hybridization, the probes of the present invention can usefully have detectable labels. Nucleic acid labels are well known in the art, and include, inter alia, radioactive labels, such as ³H, ³²P, ³³P, ³⁵S, ¹²⁵I, ¹³¹I; fluorescent labels, such as Cy3, Cy5, Cy5.5, Cy7, SYBR®

Green and other labels described in Haugland,

Handbook of Fluorescent Probes and Research Chemicals, 7th
ed., Molecular Probes Inc., Eugene, OR (2000), or
fluorescence resonance energy transfer tandem conjugates
thereof; labels suitable for chemiluminescent and/or
enhanced chemiluminescent detection; labels suitable for

ESR and NMR detection; and labels that include one member

of a specific binding pair, such as biotin, digoxigenin, or the like.

The probes, either in quantity sufficient for hybridization or sufficient for amplification, can be provided in individual vials or containers.

Alternatively, such probes can usefully be packaged as a plurality of such individual genome-derived single exon probes.

When provided as a collection of plural

individual probes, the probes are typically made available
in amplifiable form in a spatially-addressable ordered set,
typically one per well of a microtiter dish. Although a 96
well microtiter plate can be used, greater efficiency is
obtained using higher density arrays.

If, as earlier mentioned, the ORF-specific
5' primers used for genomic amplification had a first
common sequence added thereto, and the ORF-specific 3'
primers used for genomic amplification had a second,
different, common sequence added thereto, a single set of
5' and 3' primers can be used to amplify all of the probes
from the amplifiable ordered set.

Such collections of genome-derived single exon probes can usefully include a plurality of probes chosen for the common attribute of expression in the human brain.

In such defined subsets, typically at least 50, 60, 75, 80, 85, 90 or 95% or more of the probes will be chosen by their expression in the defined tissue or cell type.

The single exon probes of the present invention,

30 as well as fragments of the single exon probes comprising
selectively hybridizable portions of the probe ORF, can be
used to obtain the full length cDNA that includes the ORF
by (i) screening of cDNA libraries; (ii) rapid
amplification of cDNA ends ("RACE"); or (iii) other

35 conventional means, as are described, inter alia, in

Ausubel et al. and Maniatis et al.

It is another aspect of the present invention to provide genome-derived single exon nucleic acid microarrays useful for gene expression analysis, where the term

5 "microarray" has the meaning given in the definitional section of this description, supra.

The invention particularly provides genomederived single-exon nucleic acid microarrays comprising a plurality of probes known to be expressed in human brain. In preferred embodiments, the present invention provides human genome-derived single exon microarrays comprising a plurality of probes drawn from the group consisting of SEQ ID NOS.: 1 - 12,821.

When used for gene expression analysis, the 15 genome-derived single exon microarrays provide greater physical informational density than do the genome-derived single exon microarrays that have lower percentages of probes known to be expressed commonly in the tested tissue. At a fixed probe density, for example, a given microarray 20 surface area of the defined subset genome-derived single exon microarray can yield a greater number of expression measurements. Alternatively, at a given probe density, the same number of expression measurements can be obtained from a smaller substrate surface area. Alternatively, at a 25 fixed probe density and fixed surface area, probes can be provided redundantly, providing greater reliability in signal measurement for any given probe. Furthermore, with a higher percentage of probes known to be expressed in the assayed tissue, the dynamic range of the detection means 30 can be adjusted to reveal finer levels discrimination among the levels of expression.

Although particularly described with respect to their utility as probes of gene expression, particularly as probes to be included on a genome-derived single exon

35 microarray, each of the nucleic acids having SEQ ID NOS.: 1

- 12,821 contains an open-reading frame, set forth respectively in SEQ ID NOS.: 12,822 - 25,434, that encodes a protein domain. Thus, each of SEQ ID NOS. 1 - 12,821 can be used, or that portion thereof in SEQ ID NOS. 12,822 -5 25,434 used, to express a protein domain by standard in vitro recombinant techniques. See Ausubel et al. and Maniatis et al.

Additionally, kits are available commercially that readily permit such nucleic acids to be expressed as 10 protein in bacterial cells, insect cells, or mammalian cells, as desired (e.g., HAT Protein Expression & Purification System, ClonTech Laboratories, Palo Alto, CA; Adeno-X™ Expression System, ClonTech Laboratories, Palo Alto, CA; Protein Fusion & Purification (pMAL $^{\text{TM}}$) System, New 15 England Biolabs, Beverley, MA)

Furthermore, shorter peptides can be chemically synthesized using commercial peptide synthesizing equipment and well known techniques. Procedures are described, inter alia, in Chan et al. (eds.), Fmoc Solid Phase Peptide 20 Synthesis: A Practical Approach (Practical Approach Series, (Paper)), Oxford Univ. Press (March 2000) (ISBN: 0199637245); Jones, Amino Acid and Peptide Synthesis (Oxford Chemistry Primers, No 7) , Oxford Univ. Press (August 1992) (ISBN: 0198556683); and Bodanszky, Principles 25 of Peptide Synthesis (Springer Laboratory), Springer Verlag (December 1993) (ISBN: 0387564314).

It is, therefore, another aspect of the invention to provide peptides comprising an amino acid sequence translated from SEQ ID NOS.: 12,822 - 25,434. Such amino 30 acid sequences are set out in SEQ ID NOS: 25,435 - 37,811. Any such recombinantly-expressed or synthesized peptide of at least 8, and preferably at least about 15, amino acids, can be conjugated to a carrier protein and used to generate antibody that recognizes the peptide. Thus, it is a 35 further aspect of the invention to provide peptides that

have at least 8, preferably at least 15, consecutive amino acids.

The following examples are offered by way of illustration and not by way of limitation.

EXAMPLE 1

Preparation of Single Exon Microarrays from ORFs Predicted in Human Genomic Sequence

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Bioinformatics Results

All human BAC sequences in fewer than 10 pieces that had been accessioned in a five month period immediately preceding this study were downloaded from

15 GenBank. This corresponds to ~2200 clones, totaling ~350 MB of sequence, or approximately 10% of the human genome.

After masking repetitive elements using the program CROSS_MATCH, the sequence was analyzed for open reading frames using three separate gene finding programs.

The three programs predict genes using independent algorithmic methods developed on independent training sets:

GRAIL uses a neural network, GENEFINDER uses a hidden Markoff model, and DICTION, a program proprietary to Genetics Institute, operates according to a different

25 heuristic. The results of all three programs were used to create a prediction matrix across the segment of genomic DNA.

The three gene finding programs yielded a range of results. GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed. GENEFINDER was second, calling 1%, and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

The consensus data were as follows. GRAIL and 35 GENEFINDER agreed on 0.7% of genomic sequence, GRAIL and

DICTION agreed on 0.5% of genomic sequence, and the three programs together agreed on 0.25% of the data analyzed. That is, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding region.

ORFs predicted by any two of the three programs ("consensus ORFs") were assorted into "gene bins" using two criteria: (1) any 7 consecutive exons within a 25 kb window were placed together in a bin as likely contributing to a single gene, and (2) all ORFs within a 25 kb window were placed together in a bin as likely contributing to a single gene if fewer than 7 exons were found within the 25 kb window.

15 PCR

The largest ORF from each gene bin that did not span repetitive sequence was then chosen for amplification, as were all consensus ORFs longer than 500 bp. This method approximated one exon per gene; however, a number of genes were found to be represented by multiple elements.

Previously, we had determined that DNA fragments fewer than 250 bp in length do not bind well to the aminomodified glass surface of the slides used as support substrate for construction of microarrays; therefore, amplicons were designed in the present experiments to approximate 500 bp in length.

Accordingly, after selecting the largest ORF per gene bin, a 500 bp fragment of sequence centered on the ORF was passed to the primer picking software, PRIMER3

30 (available online for use at http://www-genome.wi.mit.edu/cgi-bin/primer/). A first additional sequence was commonly added to each ORF-unique 5' primer, and a second, different, additional sequence was commonly added to each ORF-unique 3' primer, to permit

35 subsequent reamplification of the amplicon using a single

set of "universal" 5' and 3' primers, thus immortalizing the amplicon. The addition of universal priming sequences also facilitates sequence verification, and can be used to add a cloning site should some ORFs be found to warrant 5 further study.

The ORFs were then PCR amplified from genomic DNA, verified on agarose gels, and sequenced using the universal primers to validate the identity of the amplicon to be spotted in the microarray.

10

Primers were supplied by Operon Technologies (Alameda, CA). PCR amplification was performed by standard techniques using human genomic DNA (Clontech, Palo Alto, CA) as template. Each PCR product was verified by SYBR® green (Molecular Probes, Inc., Eugene, OR) staining of 15 agarose gels, with subsequent imaging by Fluorimager (Molecular Dynamics, Inc., Sunnyvale, CA). PCR amplification was classified as successful if a single band appeared.

The success rate for amplifying ORFs of interest 20 directly from genomic DNA using PCR was approximately 75%. FIG. 5 graphs the distribution of predicted ORF (exon) length and distribution of amplified PCR products, with ORF length shown in red and PCR product length shown in blue (which may appear black in the figure). Although the range 25 of ORF sizes is readily seen to extend to beyond 900 bp, the mean predicted exon size was only 229 bp, with a median size of 150 bp (n=9498). With an average amplicon size of 475 ± 25 bp, approximately 50% of the average PCR amplification product contained predicted coding region, 30 with the remaining 50% of the amplicon containing either intron, intergenic sequence, or both.

Using a strategy predicated on amplifying about 500 bp, it was found that long exons had a higher PCR failure rate. To address this, the bioinformatics process 35 was adjusted to amplify 1000, 1500 or 2000 bp fragments

from exons larger than 500 bp. This improved the rate of successful amplification of exons exceeding 500 bp, constituting about 9.2% of the exons predicted by the gene finding algorithms.

Approximately 75% of the probes disposed on the array (90% of those that successfully PCR amplified) were sequence-verified by sequencing in both the forward and reverse direction using MegaBACE sequencer (Molecular Dynamics, Inc., Sunnyvale, CA), universal primers, and standard protocols.

Some genomic clones (BACs) yielded very poor PCR and sequencing results. The reasons for this are unclear, but may be related to the quality of early draft sequence or the inclusion of vector and host contamination in some submitted sequence data.

Although the intronic and intergenic material flanking coding regions could theoretically interfere with hybridization during microarray experiments, subsequent empirical results demonstrated that differential expression ratios were not significantly affected by the presence of noncoding sequence. The variation in exon size was similarly found not to affect differential expression ratios significantly; however, variation in exon size was observed to affect the absolute signal intensity (data not shown).

The 350 MB of genomic DNA was, by the above-described process, reduced to 9750 discrete probes, which were spotted in duplicate onto glass slides using commercially available instrumentation (MicroArray GenII Spotter and/or MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Each slide additionally included either 16 or 32 E. coli genes, the average hybridization signal of which was used as a measure of background biological noise.

Each of the probe sequences was BLASTed against

the human EST data set, the NR data set, and SwissProt GenBank (May 7, 1999 release 2.0.9).

One third of the probe sequences (as amplified) produced an exact match (BLAST Expect ("E") values less

5 than 1 e⁻¹⁰⁰) to either an EST (20% of sequences) or a known mRNA (13% of sequences). A further 22% of the probe sequences showed some homology to a known EST or mRNA (BLAST E values from 1 e⁻⁵ to 1 e⁻⁹⁹). The remaining 45% of the probe sequences showed no significant sequence homology to any expressed, or potentially expressed, sequences present in public databases.

All of the probe sequences (as amplified) were then analyzed for protein similarities with the SwissProt database using BLASTX, Gish et al., Nature Genet. 3:266

(1993). The predicted functional breakdowns of the 2/3 of probes identical or homologous to known sequences are presented in Table 1.

Table 1

Function	of Predic	ted ORFs As	Deduced From Comparative
Sequence	Analysis		
Total	V6 chip	V7 chip	Function Predicted from
			Comparative Sequence
			Analysis
211	96	115	Receptor
120	43	77	Zinc Finger
30	11	19	Homeobox
25	9	16	Transcription Factor
17	11	7	Transcription
118	57	61	Structural
95	39	56	Kinase
36	18	18	Phosphatase
83	31	52	Ribosomal

45	19	26	Transport
21	17	14	Growth Factor
17	12	5	Cytochrome
50	33	17	Channel

As can be seen, the two most common types of genes were transcription factors and receptors, making up 2.2% and 1.8% of the arrayed elements, respectively.

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EXAMPLE 2

Gene Expression Measurements From Genome-Derived Single Exon Microarrays

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The two genome-derived single exon microarrays prepared according to Example 1 were hybridized in a series of simultaneous two-color fluorescence experiments to (1)

15 Cy3-labeled cDNA synthesized from message drawn individually from each of brain, heart, liver, fetal liver, placenta, lung, bone marrow, HeLa, BT 474, or HBL 100 cells, and (2) Cy5-labeled cDNA prepared from message pooled from all ten tissues and cell types, as a control in each of the measurements. Hybridization and scanning were carried out using standard protocols and Molecular Dynamics equipment.

Briefly, mRNA samples were bought from commercial sources (Clontech, Palo Alto, CA and Amersham Pharmacia
25 Biotech (APB)). Cy3-dCTP and Cy5-dCTP (both from APB) were incorporated during separate reverse transcriptions of 1 µg of polyA⁺ mRNA performed using 1 µg oligo(dT)12-18 primer and 2 µg random 9mer primers as follows. After heating to 70°C, the RNA:primer mixture was snap cooled on ice. After snap cooling on ice, added to the RNA to the stated final concentration was: 1X Superscript II buffer, 0.01 M DTT,

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100μM dATP, 100 μM dGTP, 100 μM dTTP, 50 μM dCTP, 50 μM Cv3-dCTP or Cv5-dCTP 50 µM, and 200 U Superscript II The reaction was incubated for 2 hours at 42°C. After 2 hours, the first strand cDNA was isolated by adding 5 1 U Ribonuclease H, and incubating for 30 minutes at 37°C. The reaction was then purified using a Qiagen PCR cleanup column, increasing the number of ethanol washes to 5. Probe was eluted using 10 mM Tris pH 8.5.

Using a spectrophotometer, probes were measured 10 for dye incorporation. Volumes of both Cy3 and Cy5 cDNA corresponding to 50 pmoles of each dye were then dried in a Speedvac, resuspended in 30 µl hybridization solution containing 50% formamide, 5X SSC, 0.2 µg/µl poly(dA), 0.2 ug/ul human cot1 DNA, and 0.5 % SDS.

15

Hybridizations were carried out under a coverslip, with the array placed in a humid oven at 42°C overnight. Before scanning, slides were washed in 1X SSC, 0.2% SDS at 55°C for 5 minutes, followed by 0.1X SSC, 0.2% SDS, at 55°C for 20 minutes. Slides were briefly dipped in 20 water and dried thoroughly under a gentle stream of nitrogen.

Slides were scanned using a Molecular Dynamics Gen3 scanner, as described. Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing 25 Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

Although the use of pooled cDNA as a reference permitted the survey of a large number of tissues, it attenuates the measurement of relative gene expression, 30 since every highly expressed gene in the tissue/cell typespecific fluorescence channel will be present to a level of at least 10% in the control channel. Because of this fact, both signal and expression ratios (the latter hereinafter, "expression" or "relative expression") for each probe were 35 normalized using the average ratio or average signal,

respectively, as measured across the whole slide.

Data were accepted for further analysis only when signal was at least three times greater than biological noise, the latter defined by the average signal produced by the E. coli control genes.

The relative expression signal for these probes was then plotted as function of tissue or cell type, and is presented in FIG. 6.

FIG. 6 shows the distribution of expression

10 across a panel of ten tissues. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), and expressed in all tissues tested ("10").

Of 9999 arrayed elements on the two microarrays (including positive and negative controls and "failed" products), 2353 (51%) were expressed in at least one tissue or cell type. Of the gene elements showing significant signal — where expression was scored as "significant" if the normalized Cy3 signal was greater than 1, representing signal 5-fold over biological noise (0.2) — 39% (991) were expressed in all 10 tissues. The next most common class (15%) consisted of gene elements expressed in only a single tissue.

The genes expressed in a single tissue were further analyzed, and the results of the analyses are compiled in FIG. 7.

FIG. 7A is a matrix presenting the expression of all verified sequences that showed expression greater than 30 3 in at least one tissue. Each clone is represented by a column in the matrix. Each of the 10 tissues assayed is represented by a separate row in the matrix, and relative expression of a clone in that tissue is indicated at the respective node by intensity of green shading, with the intensity legend shown in panel B. The top row of the

matrix ("EST Hit") contains "bioinformatic" rather than
 "physical" expression data - that is, presents the results
 returned by query of EST, NR and SwissProt databases using
 the probe sequence. The legend for "bioinformatic
 expression" (i.e., degree of homology returned) is

presented in panel C. Briefly, white is known, black is novel, with gray depicting nonidentical with significant homology (white: E values < 1e-100; gray: E values from 1e-05 to 1e-99; black: E values > 1e-05).

As FIG. 7 readily shows, heart and brain were demonstrated to have the greatest numbers of genes that were shown to be uniquely expressed in the respective tissue. In brain, 200 uniquely expressed genes were identified; in heart, 150. The remaining tissues gave the following figures for uniquely expressed genes: liver, 100; lung, 70; fetal liver, 150; bone marrow, 75; placenta, 100; HeLa, 50; HBL, 100; and BT474, 50.

"novel" genes among those that were up-regulated in only
20 one tissue, as compared with those that were down-regulated
in only one tissue. In fact, it was found that ORFs whose
expression was measurable in only a single of the tested
tissues were represented in sequencing databases at a rate
of only 11%, whereas 36% of the ORFs whose expression was
25 measurable in 9 of the tissues were present in public
databases. As for those ORFs expressed in all ten tissues,
fully 45% were present in existing expressed sequence
databases. These results are not unexpected, since genes
expressed in a greater number of tissues have a higher
30 likelihood of being, and thus of having been, discovered by
EST approaches.

Comparison of Signal from Known and Unknown Genes

The normalized signal of the genes found to have high homology to genes present in the GenBank human EST

database were compared to the normalized signal of those genes not found in the GenBank human EST database. The data are shown in FIG. 8.

FIG. 8 shows the normalized Cy3 signal intensity

for all sequence-verified products with a BLAST Expect
("E") value of greater than 1e-30 (designated "unknown")
upon query of existing EST, NR and SwissProt databases, and
shows in blue the normalized Cy3 signal intensity for all
sequence-verified products with a BLAST Expect value of

less than 1e-30 ("known"). Note that biological background
noise has an averaged normalized Cy3 signal intensity of
0.2.

As expected, the most highly expressed of the ORFs were "known" genes. This is not surprising, since very high signal intensity correlates with very commonly-expressed genes, which have a higher likelihood of being found by EST sequence.

However, a significant point is that a large number of even the high expressers were "unknown". Since the genomic approach used to identify genes and to confirm their expression does not bias exons toward either the 3' or 5' end of a gene, many of these high expression genes will not have been detected in an end-sequenced cDNA library.

The significant point is that presence of the gene in an EST database is *not* a prerequisite for incorporation into a genome-derived microarray, and further, that arraying such "unknown" exons can help to assign function to as-yet undiscovered genes.

30

Verification of Gene Expression

To ascertain the validity of the approach described above to identify genes from raw genomic sequence, expression of two of the probes was assayed using reverse transcriptase polymerase chain reaction (RT PCR)

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and northern blot analysis.

10

Two microarray probes were selected on the basis of exon size, prior sequencing success, and tissue-specific gene expression patterns as measured by the microarray 5 experiments. The primers originally used to amplify the two respective ORFs from genomic DNA were used in RT PCR against a panel of tissue-specific cDNAs (Rapid-Scan gene expression panel 24 human cDNAs) (OriGene Technologies, Inc., Rockville, MD).

Sequence AL079300 1 was shown by microarray hybridization to be present in cardiac tissue, and sequence AL031734 1 was shown by microarray experiment to be present in placental tissue (data not shown). RT-PCR on these two sequences confirmed the tissue-specific gene expression as 15 measured by microarrays, as ascertained by the presence of a correctly sized PCR product from the respective tissue type cDNAs.

Clearly, all microarray results cannot, and indeed should not, be confirmed by independent assay 20 methods, or the high throughput, highly parallel advantages of microarray hybridization assays will be lost. However, in addition to the two RT-PCR results presented above, the observation that 1/3 of the arrayed genes exist in expression databases provides powerful confirmation of the 25 power of our methodology - which combines bioinformatic prediction with expression confirmation using genomederived single exon microarrays - to identify novel genes from raw genomic data.

To verify that the approach further provides 30 correct characterization of the expression patterns of the identified genes, a detailed analysis was performed of the microarrayed sequences that showed high signal in brain.

For this latter analysis, sequences that showed high (normalized) signal in brain, but which showed very 35 low (normalized) signal (less than 0.5, determined to be

biological noise) in all other tissues, were further studied. There were 82 sequences that fit these criteria, approximately 2% of the arrayed elements. The 10 sequences showing the highest signal in brain in microarray

5 hybridizations are detailed in Table 2, along with assigned function, if known or reasonably predicted.

Table 2

I	unction	of the Mo	st Highly	
Expressed 0				n
	~		-, Diui	••
Microarray	Normal	Expressi	Homology	Gene Function
Sequence	ized	on Ratio	to EST	as described by
Name	Signal		present	
			in	
			GenBank	
AP000217-1	5.2	+7.7	High	S-100 protein,
				b-chain, Ca ²⁺
				binding protein
				expressed in
				central nervous
				system
AP000047-1	2.3		High	Unknown
				Function
AC006548-9	1.7		High	Similar to
				mouse membrane
				glyco-protein
				M6, expressed
•				in central
	İ			nervous system

AC007245-5	1.5	<u> </u>	High	Similar to
			911	amphiphysin, a
			'	synaptic
				vesicle-
,				associated
				protein. Ref 21
L44140-4	1.2	+2.0	rid ah	Endothelial
144140-4	1.2	+2.0	High	
				actin-binding
				protein found
				in nonmuscle
				filamin
AC004689-9	1.2	+3.5	High	Protein
				Phosphatase
				PP2A, neuronal/
				downregulates
				activated
1				protein kinases
AL031657-1	1.2	+3.0	High	Unknown
				function/
				Contains the
				anhyrin motif,
				a common
				protein
				sequence motif
AC009266-2	1.1	+3.7	Low	Low homology to
				the
				Synaptotagmin I
				protein in
				rat/present at
				low levels
				throughout rat
				brain
AP000086-1	1.0	+2.7	Low	Unknown, very
				poor homology
				to collagen
1	l			
		88		

Protein Phosphatase
Phosphatase
PP2A, neuronal/
downregulates
activated
protein kinases

Of the ten sequences studied by these latter confirmatory approaches, eight were previously known. Of these eight, six had previously been reported to be important in the central nervous system or brain. The exon giving the highest signal (AP00217-1) was found to be the gene encoding an S100B Ca²⁺ binding protein, reported in the literature to be highly and uniquely expressed in the central nervous system. Heizmann, Neurochem. Res. 9:1097 (1997).

A number of the brain-specific probe sequences (including AC006548-9, AC009266-2) did not have homology to any known human cDNAs in GenBank but did show homology to rat and mouse cDNAs. Sequences AC004689-9 and AC004689-3

15 were both found to be phosphatases present in neurons (Millward et al., Trends Biochem. Sci. 24(5):186-191 (1999)). Two microarray sequences, AP000047-1 and AP000086-1 have unknown function, with AP000086-1 being absent from GenBank. Functionality can now be narrowed down to a role in the central nervous system for both of these genes, showing the power of designing microarrays in this fashion.

Next, the function of the chip sequences with the highest (normalized) signal intensity in brain, regardless of expression in other tissues, was assessed. In this latter analysis, we found expression of many more common genes, since the sequences were not limited to those expressed only in brain. For example, looking at the 20 highest signal intensity spots in brain, 4 were similar to

tubulin (AC00807905; AF146191-2; AC007664-4; AF14191-2), 2 were similar to actin (AL035701-2; AL034402-1), and 6 were found to be homologous to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (AL035604-1; Z86090-1; AC006064-L, AC006064-K; AC035604-3; AC006064-L). These genes are often used as controls or housekeeping genes in microarray experiments of all types.

Other interesting genes highly expressed in brain were a ferritin heavy chain protein, which is reported in the literature to be found in brain and liver (Joshi et al., J. Neurol. Sci. 134(Suppl):52-56 (1995)), a result duplicated with the array. Other highly expressed chip sequences included a translation elongation factor 10 (AC007564-4), a DEAD-box homolog (AL023804-4), and a Y-chromosome RNA-binding motif (Chai et al., Genomics 49(2):283-89 (1998)) (AC007320-3). A low homology analog (AP00123-1/2) to a gene, DSCR1, thought to be involved in trisomy 21 (Down's syndrome), showed high expression in both brain and heart, in agreement with the literature (Fuentes et al., Mol. Genet. 4(10):1935-44 (1995)).

As a further validation of the approach, we selected the BAC AC006064 to be included on the array. This BAC was known to contain the GAPDH gene, and thus could be used as a control for the ORF selection process.

25 The gene finding and exon selection algorithms resulted in choosing 25 exons from BAC AC006064 for spotting onto the array, of which four were drawn from the GAPDH gene. Table 3 shows the comparison of the average expression ratio for the 4 exons from BAC006064 compared with the average expression ratio for 5 different dilutions of a commercially available GAPDH cDNA (Clontech).

Table 3

Comp	arigon of B	
tissue, of GAPI	arison of Expression DH	Ratio, for each
	AC006064 (n = 4)	Control (n = 5)
Bone Marrow	-1.81 ± 0.11	-1.85 ± 0.08
Brain	-1.41 ± 0.11	-1.17 ± 0.05
BT474	1.85 ± 0.09	1.66 ± 0.12
Fetal Liver	-1.62 ± 0.07	-1.41 ± 0.05
HBL100	1.32 ± 0.05	2.64 ± 0.12
Heart	1.16 ± 0.09	1.56 ± 0.10
HeLa	1.11 ±0.06	1.30 ± 0.15
Liver	-1.62 ± 0.22	-2.07 ±
Lung	-4.95 ± 0.93	-3.75 ± 0.21
Placenta	-3.56 ± 0.25	-3.52 ± 0.43

Each tissue shows excellent agreement between the experimentally chosen exons and the control, again

demonstrating the validity of the present exon mining approach. In addition, the data also show the variability of expression of GAPDH within tissues, calling into question its classification as a housekeeping gene and utility as a housekeeping control in microarray experiments.

EXAMPLE 3

Representation of Sequence and Expression Data as a "Mondrian"

15

For each genomic clone processed for microarray as above-described, a plethora of information was accumulated, including full clone sequence, probe sequence within the clone, results of each of the three gene finding programs, EST information associated with the probe

sequences, and microarray signal and expression for multiple tissues, challenging our ability to display the information.

Accordingly, we devised a new tool for visual display of the sequence with its attendant annotation which, in deference to its visual similarity to the paintings of Piet Mondrian, is hereinafter termed a "Mondrian". FIGS. 3 and 4 present the key to the information presented on a Mondrian.

10 FIG. 9 presents a Mondrian of BAC AC008172 (bases 25,000 to 130,000 shown), containing the carbamyl phosphate synthetase gene (AF154830.1). Purple background within the region shown as field 81 in FIG. 3 indicates all 37 known exons for this gene.

As can be seen, GRAIL II successfully identified 27 of the known exons (73%), GENEFINDER successfully identified 37 of the known exons (100%), while DICTION identified 7 of the known exons (19%).

Seven of the predicted exons were selected for physical assay, of which 5 successfully amplified by PCR and were sequenced. These five exons were all found to be from the same gene, the carbamyl phosphate synthetase gene (AF154830.1).

The five exons were arrayed, and gene expression
25 measured across 10 tissues. As is readily seen in the
Mondrian, the five chip sequences on the array show
identical expression patterns, elegantly demonstrating the
reproducibility of the system.

FIG. 10 is a Mondrian of BAC AL049839. We

selected 12 exons from this BAC, of which 10 successfully sequenced, which were found to form between 5 and 6 genes. Interestingly, 4 of the genes on this BAC are protease inhibitors. Again, these data elegantly show that exons selected from the same gene show the same expression patterns, depicted below the red line. From this figure,

it is clear that our ability to find known genes is very
good. A novel gene is also found from 86.6 kb to 88.6 kb,
upon which all the exon finding programs agree. We are
confident we have two exons from a single gene since they
show the same expression patterns and the exons are
proximal to each other. Backgrounds in the following
colors indicate a known gene (top to bottom):
red = kallistatin protease inhibitor (P29622);
purple = plasma serine protease inhibitor (P05154);
turquoise = α1 anti-chymotrypsin (P01011); mauve = 40S
ribosomal protein (P08865). Note that chip sequence 8 and
12 did not sequence verify.

15 EXAMPLE 4

Genome-Derived Single Exon Probes Useful For Measuring Human Gene Expression

The protocols set forth in Examples 1 and 2,

20 supra, were applied to additional human genomic sequence as

it became newly available in GenBank to identify unique

exons in the human genome that could be shown to be

expressed at significant levels in brain tissue.

These unique exons are within longer probe

25 sequences. Each probe was completely sequenced on both strands prior to its use on a genome-derived single exon microarray; sequencing confirms the exact chemical structure of each probe. An added benefit of sequencing is that it placed us in possession of a set of single base
30 incremented fragments of the sequenced nucleic acid, starting from the sequencing primer 3' OH. (Since the single exon probes were first obtained by PCR amplification from genomic DNA, we were of course additionally in possession of an even larger set of single base incremented fragments of each of the 12,821 single exon probes, each

fragment corresponding to an extension product from one of the two amplification primers.)

The structures of the 12,821 unique single exon probes are clearly presented in the Sequence Listing as SEQ ID Nos.: 1 - 12,821. The 16 nt 5' primer sequence and 16 nt 3' primer sequence present on the amplicon are not included in the sequence listing. The sequences of the exons present within each of these probes is presented in the Sequence Listing as SEQ ID Nos.: 12,822 - 25,434, respectively. It will be noted that some amplicons have more than one exon, some exons are contained in more than one amplicon.

As detailed in Example 2, expression was
demonstrated by disposing the amplicons as single exon
probes on nucleic acid microarrays and then performing twocolor fluorescent hybridization analysis; significant
expression is based on a statistical confidence that the
signal is significantly greater than negative biological
control spots. The negative biological control is formed
from spotted DNA sequences from a different species. Here,
32 sequences from E.Coli were spotted in duplicate to give
a total of 64 spots.

For each hybridisation (each slide, each colour) the median value of the signal from all of the spots is determined. The normalised signal value is the arithmetic mean of the signal from duplicate spots divided by the population median.

Control spots are eliminated if there is more that a five-fold difference between each one of the duplicate spots raw signals.

The median of the signal from the remaining control spots is calculated and all subsequent calculations are done with normalised signals.

Control spots having a signal of greater than 35 median + 2.4 (the value 2.4 is roughly 12 times the

observed standard deviation of control spot populations) are eliminated. Spots with such high signals are considered to be "outliers".

The mean and standard deviation of the modified control spot populations are calculated.

The mean + 3x the standard deviation (mean + (3*SD)) is used as the signal threshold qualifier for that particular hybridisation. Thus, individual thresholds are determined for each channel and each hybridisation.

This means that, assuming that the data is distributed normally, there is a 99% confidence that any signal exceeding the threshold is significant.

The probes and their expression data are presented in Table 4, set forth respectively in Example 5.

Example 5 presents the subset of probes that is significantly expressed in the human heart and thus presents the subset of probes that was recognized to be useful for measuring expression of their cognate genes in human brain tissue.

The sequence of each of the exon probes identified by SEQ ID NOS.: 12,822 - 25,434 was individually used as a BLAST (or, for SWISSPROT, BLASTX) query to identify the most similar sequence in each of dbEST, SwissProt (BLASTX), and NR divisions of GenBank. Because the query sequences are themselves derived from genomic sequence in GenBank, only nongenomic hits from NR were scored.

The smallest in value of the BLAST (or BLASTX) expect ("E") scores for each query sequence across the

three database divisions was used as a measure of the
"expression novelty" of the probe's ORF. Table 4 is sorted in descending order based on this measure, reported as
"Most Similar (top) Hit BLAST E Value". Those sequences for which no "Hit E Value" is listed are those exons which were

found to have no similar sequences.

As sorted, Table 4 thus lists its respective probes (by "AMPLICON SEQ ID NO.:" and additionally by the SEQ ID NO:. of the exon contained within the probe: "EXON SEQ ID NO.:") from least similar to sequences known to be expressed (i.e., highest BLAST E value), at the beginning of the table, to most similar to sequences known to be expressed (i.e., lowest BLAST E value), at the bottom of the table.

Table 4 further provides, for each listed probe,

the accession number of the database sequence that yielded
the "Most Similar (top) Hit BLAST E Value", along with the
name of the database in which the database sequence is
found ("Top Hit Database Source").

Table 4 further provides SEQ ID NOS.

15 corresponding to the predicted amino acid sequences where they have been determined for the probe and exon nucleotide sequences. These are set out as PEPTIDE SEQ ID NOS.:. The peptide sequences for a given exon are predicted as follows: Since each chip exon is a consensus sequence drawn 20 from predictions from various exon finding programs (i.e. Grail, GeneFinder and GenScan), the multiple initial ORFs are first determined in a uniform way according to each prediction. In particular, the reading frame for predicting the first amino acid in the peptide sequence always starts 25 with the first base of any codon and ends with the last base of non-termination codon. Next, for each strand of the exon, initial ORFs are merged into one or more final ORFs in an exhaustive process based on the following criteria: 1) the merging ORFs must be overlapping, and 2) the merging 30 ORFs must be in the same frame.

The Sequence Listing, which is a superset of all of the data presented in Table 4, further includes, for each probe, the most similar hit, with accession number and BLAST E value, from the each of the three queried databases.

Table 4 further lists, for each probe, a portion. of the descriptor for the top hit ("Top Hit Descriptor") as provided in the sequence database. For those ORFs that are similar in sequence, but nonidentical to known sequences (e.g., those with BLAST E values between about 1e-05 and 1e-100), the descriptor reveals the likely function of the protein encoded by the probe's ORF.

Using BLAST E value cutoffs of 1e-05 (i.e., 1 x 10⁻⁵) and 1e-100 (i.e., 1 x 10⁻¹⁰⁰) as evidence of similarity to sequences known to be expressed is of course arbitrary: in Example 2, supra, a BLAST E value of 1e-30 was used as the boundary when only two classes were to be defined for analysis (unknown, >1e-30; known <1e-30) (see also FIG. 8). Furthermore, even when the "Most Similar (Top) Hit BLAST E Value" is low, e.g., less than about 1e-100 - which is probative evidence that the query sequence has previously been shown to be expressed - the top hit is highly unlikely exactly to match the probe sequence.

First, such expression entries typically will not 20 have the intronic and/or intergenic sequence present within the single exon probes listed in the Table. Second, even the ORF itself is unlikely in such cases to be present identically in the databases, since most of the EST and mRNA clones in existing databases include multiple exons, 25 without any indication of the location of exon boundaries.

As noted, the data presented in Table 4 represent a proper subset of the data present within the attached sequence listing. For each amplicon probe (SEQ ID NOs.: 1 - 12,821) and probe exon (SEQ ID NOs.: 12,822 - 25,434, 30 respectively), the sequence listing further provides, through iterated annotation fields <220> and <223>:

(a) the accession number of the BAC from which the sequence was derived ("MAP TO"), thus providing a link to the chromosomal map location and other information about the genomic milieu of the probe sequence;

(b) the most similar sequence provided by BLAST query of the EST database, with accession number and BLAST E value for the "hit";

- (c) the most similar sequence provided by BLAST 5 query of the GenBank NR database, with accession number and BLAST E value for the "hit"; and
 - (d) the most similar sequence provided by BLASTX query of the SWISSPROT database, with accession number and BLAST E value for the "hit".

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EXAMPLE 5

Genome-Derived Single Exon Probes Useful For Measuring Expression of Genes in Human Brain

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 $\overline{\text{Table 4}}$ (536 pages) presents expression, homology, and functional information for the genome-derived single exon probes that are expressed significantly in human brain.

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CLAIMS

A spatially-addressable set of single exon nucleic acid probes for measuring gene expression in a sample derived
 from human brain comprising a plurality single exon nucleic probes, said probes comprising any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 12,821 or a complementary sequence, or a portion of such a sequence.

- 10 2. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably amplifiable.
- 3. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably isolatable from said plurality.
- 4. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 3 wherein said probes comprise any one of the nucleotide sequences set out in SEQ ID NOS.: 12,822 - 25,434.
- 5. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 4, wherein each of said plurality of probes is amplifiable using at least one common primer.
- 6. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 5 wherein the set comprises between 50 20,000 single exon nucleic acid probes.
- 7. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 6, wherein the

average length of the single exon nucleic acid probes is between 200 and 500 bp.

- 8. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 7, wherein at least 50% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.
- 9. A spatially-addressable set of single exon nucleic acid 10 probes as claimed in any of claims 1 to 8, wherein at least 50% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.
- 10. A spatially-addressable set of single exon nucleic acid 15 probes as claimed in any of claims 1 - 9 characterised in that said set of probes is addressably disposed upon a substrate.
- 11. A spatially-addressable set of single exon nucleic acid 20 probes as claimed in claim 10 wherein said substrate is selected from glass, amorphous silicon, crystalline silicon and plastic.
- 12. A microarray comprising a spatially addressable set of25 single exon nucleic acid probes as claimed in any of claims1 11.
- 13. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 12,821 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid molecule expressed in the human brain.

14. A single exon nucleic acid probe as claimed in claim 13 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 12,822 - 25,434 or a complementary sequence or a fragment thereof.

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- 15. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 25,435 37,811, or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human brain.
- 16. A single exon nucleic acid probe as claimed in any one of claims 13 to 15 wherein said single exon nucleic acid probe comprises between 15 and 25 contiguous nucleotides of said SEQ ID NO.
- 17. A single exon nucleic acid probe as claimed in any one of claims 13 to 15, wherein said probe is between 3 25 kb in length.
 - 18. A single exon nucleic acid probe as claimed in any one of claims 13 17, wherein said probe is DNA, RNA or PNA.

25

- 19. A single exon nucleic acid probe as claimed in any one of claims 13 18, wherein said probe is detectably labeled.
- 30 20. A single exon nucleic acid probe as claimed in any one of claims 13 19, wherein said probe lacks prokaryotic and bacteriophage vector sequence.
- 21. A single exon nucleic acid probe as claimed in any one of claims 13 20, wherein said probe lacks homopolymeric

stretches of A or T.

22. A method of measuring gene expression in a sample derived from human brain, comprising:

contacting the microarray of claim 12, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human brain; and then

measuring the label detectably bound to each probe of said microarray.

23. A method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are derived from mRNA from the brain of said eukaryote, said probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a microarray according to claim 12, and said fragment is selectively hybridizable at high stringency.

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24. A method of assigning exons to a single gene, comprising:

identifying a plurality of exons from genomic sequence according to the method of claim 23; and then

measuring the expression of each of said exons in a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe with said exon,

35 wherein a common pattern of expression of said exons in

said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

- 25. A nucleic acid sequence as set out in any of SEQ ID 5 NOs: 1 25,434 which encodes a peptide.
 - 26. A peptide encoded by a sequence as set out in any of SEQ ID Nos: 1 25,434.
- 10 27. A peptide comprising a sequence as set out in any of SEQ ID NOs: 25,435 37,811.

Page 1 of 536 Table 4 Single Exon Probes Expressed in Brain

Single Exon Probes Expressed in Brain	Top Hit Descriptor																																		
jie Exon Prope	Top Hit Database Source																																		
ŽIIO	Top Hit Acession No.																																		
	Most Similar (Top) Hit BLAST E Value																																		
	Expression Signal	6.47	15.92	2.15	10.88	1.22	1.22	3.19	6.1	3.31	1.44	6.78	1.44	2.14	2.7	2.91	1	-	2.83	1.42	12.04	1	1.67	1.03	1.52	6.4	26.0	0.97	1.07	0.78	0.99	5.38	1.3	1.57	1.67
	ORF SEQ ID NO:	[]	26308			12692				27160		27192	27331	27425	i			İ					29042			29595	29613			29734				30500	
	Exan SEQ ID NO:	13223	١.	١.	14029	14235		14355	14379	14461	14485	14492	14621	1		l I		15292				16304	16402	16678	16913	16971	16989	16989	17042	17099	17518	17708	17718		17985
	Probe SEQ ID NO:	437	889	1022	1279	1488	1488	1609	1633	1718	1743	1750	1884	1971	2162	727	2578	2578	3181	3442	3505	3549	3649	3928	4173	4230	4248	4248	4303	4361	4784	4983	4995	5176	5176

Page 2 of 536 Table 4

		·																										<u> </u>	3			B		E	1		A
Single Exon Probes Expressed in Brain	Top Hit Describer																																			00	riving suprema Log gene, partial, exons 15, 16, 17 and 18
le Exon Probes	Top Hit Database	Source	1			+			+		-		1	+	+	1			-	-			1	+	1	+	1		+	1				+		The state of the s	111011
Sing	Top Hit Acese No.											+					+	+	1	1	1	+	1	-	_		-					-	-	+	+	239028.1 INT	1
	Most Similar (Top) Hit BLAST-E	Value															1	<u> </u>				+	+	 				-	-	-					+	9.9E+00 AJ239028.1	
	Expression Signel		43	8.14	3.97	9.0	3.28	1.62	1.76	127	1	17	٦	F	1.78	1.78	0.61	4-	1.49	0.59	0.59	2.67	0.77	1.24	0.94	0.62	0.62	2.53	1.34	2.2	1.84	2.02	2.47	1.52	2.36	17.79	
	ORF SEQ ID NO:					31356	31362	31673	31699		32220	32221	32774	32775	33071	33072		33780	34214	34598	34599	35287	35515	35838	35767	36194	36195		36879		36968	37328			31006	31703	
	SEQ ID	1	18139	18308		Ш		1		19092	. 1				19994						21451				22570	22975	22975	23277	26131	23629	23701	24023	24086	24735	24916	18743	
	Probe SEQ ID NO:		5336	5510	5593	5648	. 5854	5932	5958	6322	8454	. 6454	7025	7025	7311	7311	7712	7960	8384	8759	8769	9434	9996	9782	3922	10328	10328	10582	10749	10952	11030	11332	11485	12313	1	5961	

Page 3 of 536 Table 4 Single Exon Probes Expressed in Brain

H					6		
ພ ພ ້	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1	20600	33730	1.74	9.8E+00	9.8E+00 U32716.1	NT	Haemophilus influenzae Rd section 31 of 163 of the complete genome
	22295		0.44	9.8E+00		NT	Suffolobus soffataricus 281 kb genomic DNA fragment, strain P2
	22295	35490		9.8E+00		TN	Suifolobus soffataricus 281 kb genomic DNA fragment, strain P2
	19639	32684		9.6E+00	9.6E+00 AF065630.1	NT	Gallus gallus ornithine transcarbamylase (OTC) gene, exon 1
	19639	32685		9.6E+00	AF065630.1	N	Gallus gallus ornithine transcarbarnylase (OTC) gene, exon 1
	22968		1.17		9.6E+00 AF242432.1	NT	Mus musculus Naip3 gene, exon 1; neuronal apoptosis inhibitory protein 1 (Naip1) and general transcription factor IIH polypeptide 2 (Gtf2h2) genes, complete cds
	22968	36188	1.17	9.6€+00	9.6E+00 AF242432.1	NT	Mus musculus Nalp3 gene, exon 1; neuronal apoptosis Inhibitory protein 1 (Nalp1) and general transcription factor IIH polypeptide 2 (Gtf2h2) genes, complete cds
	15381	28119	1	9.4E+00 L11433.1	111433.1	NT	Dengue virus type 3 membrane protein (prMM)/envelope glycoprotein (E) polyprotein mRNA, partial cds
	15381	28120	•	9.4E+00	9.4E+00 L11433.1	LΝ	Dengue virus type 3 membrane protein (pr/MM)/envelope glycoprotein (E) polyprotein mRNA, partial cds
	15690	28334	2.87	9.4E+00	9.4E+00 AB043785.1	IN	Mus musculus AT3 gene for antithrombin, complete cds
	20692	33820	0.91	9.3E+00	9.3E+00 AF130990.1	۲	Homo saplens ectodysplasin-A receptor protein (EDAR) gene, exons 2, 3, and 4
	21592	34733	3.06		P11210	SWISSPROT	IMMEDIATE-EARLY PROTEIN 1 (IE1) (IMMEDIATE-EARLY PHOSPHOPROTEIN PP89)
	18022	30645	2.46	9.1E+00	9.1E+00 AF095609.1	. IN	Leuciscus caphalus orientalis cytochrome b (cyt b) gene, partial cds; mitochondrial gene for mitochondrial product
	3	20646	97 0		. 1008800 4	H	Leuciscus cephalus orientalis cytochrome b (cyt.b) gene, partial ods; mitochondrial gene for mitochondrial
- 1	21997	3			ģ	SWISSPROT	RHODOPSIN
Ł	18727	31685		8.9E+00	8.8E+00 BE971806.1	EST_HUMAN	601651038R1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:3934592 3'
	19060	32041				TN	Cynops pyrrhogaster CpTbx3 premature mRNA, partial cds
	19060	32042	2.28			INT	Cynops pyrrhogaster CpTbx3 premature mRNA, partial cds
	13216	25861	2.3	8.4E+00	8.4E+00 5031804 NT	NT	Homo sapiens insulin receptor substrate 1 (IRS1) mRNA
	20426	33545	3.58	8.1E+00		IN	Zea mays mRNA for legumain-like protease (see2a)
	23791		2	8.0E+00 P41820		SWISSPROT	BREFELDIN A RESISTANCE PROTEIN
	20745		68'0	7.6E+00		IN	African swine fewer virus NP1450L gene encoding RNA polymerase largest subunit
	19931		1.9	7.5E+00	5.1	NT	Thermoplasma acidophilum complete genome; segment 3/5
	20953	34090		7.5E+00 P35441		SWISSPROT	THROMBOSPONDIN 1 PRECURSOR
	20953	34091	1.61	7.5E+0		SWISSPROT	THROMBOSPONDIN 1 PRECURSOR
	18504	31426		7.4E+0	DBF700517.1		602128876F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4285506 5
	21343	34487		7.4E+00 P04929	P04929	\Box	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
	21343	34488	2.7	7.4E+00	P04929	SWISSPROT	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR

Page 4 of 536 Table 4 Single Exon Probes Expressed in Brain

			_	$\overline{}$	_	_	_	_	_	_	_	_	_	_	_					_														
	Top Hit Descriptor	Voltare from social on the Mill OTD (A P. D. A.	Lycaper score secure in the Contract of the Co	Properties of secure from Mill. 6 i Pase (SAR2) mRNA, complete cds	ZINC FINCED DEOTEM 4 ZINC FINCED DEOTEM 2 ZINC FINC	ZING-FINGER PROTEIN 1 (ZING-FINGER HOMEODOMAIN PROTEIN 1)	Arabidansis theliam DNA of	HYPOTHETICAL 173 KDA BBOTEIN IN 1953, BUILDEN NOTE.	ARGININE KINASE (AK)	WOAR REPEAT BROTEIN IVER	60S RIBOSOMAI PROTEIN 1 4 4 3)	DNA MISMATCH REPAIR PROTEIN MITE	SKT5 PROTEIN	28070111 Source majorom to 28 Eure 1	TOTAL 1.11 Source integrational and in the supplemental and Service IMAGE:291860 6'	OUTER CAPEIN PROTEIN VP4 (HENAGGLUTININ) (OUTER LAYER PROTEIN VP4) [CONTAINS:	UNDOTHETION ATTOCKED TO THE BILL OF THE BI	CATECHOL ON ETHIC PRINCIPE OF SECTION OF THE SECTIO	CALECTION CONTRIBUTION OF THE MAN SELVEN SOUND SELVEN (S-COMT)	URIDA ATE KINASE / IN A IDENTIFY VOLUME CON CONTRACE: 4293427 5'	URIDYI ATE KINASE (111X) (IDEIDINE MONOPHOSPHATE KINASE) (UMP KINASE)	PROBABLE CATION, TRANSPORTING ATBASE ASSAURT	ENV POLYPROTEIN ICONTAINS: COAT PROTEIN OBES: COAT BROTTEIN BEST	601678435F1 NIH MGC 63 Home seniens cDNA clare IMACE 2000000 FF	Schlzophyllum corranue unknown mRNA	Mus musculus mannosidasa 2 olaha B4 (Manchata DAIA	601468031F1 NIH MGC 67 Home sealons a DNA alexanders and a least to the contract of the contra	Purcoccus harlaceli 013 general DMA 4 cons INACE:38/1303 5	Democacus radiodingers R4 scatture 1 20 2 44	Delnococcus radiodurans R1 section 1 of 2 of the complete chromosome 2	Mus musculus mixed lineage kinase 3 (Mik3) and two pore domain K+ channel subunit (Kcnk6) genes.	Homo seniere DESC1 and him (DESC1) - DA14	Mus musculus immi incolobrillo scanoscata posta i post	and a second second leading leading leading leading the second leading
	Top Hit Database Source	TZ	Ė	EST HIMAN		SWISSPROT	NT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	Г	Τ	L	Т	T	T	T	Г	SWISSPROT	Т	T	L	Γ	N.		T HUMAN	Т			2 (
	Top Hit Acession No.	+00 L12051.1	+00 1120511	+00 BE179090.1	P28166	P28166	7.1E+00 AL161595.2	P05850	P48610	022469	P35679								1	l				Г	+00 AY010901.1	6754621 NT	00 BE780163.1 E	00 AP000008.1	00 AE001862.1			1557		1
	Most Similar (Top) Hit BLAST E Value	7.2E+00	7.2E+00	7.2E+00	7.1E+00 P28166	7.1E+00 P28166	7.1E+00	7.1E+00 P05850	7.0E+00 P48610	7.0E+00 O22469	6.9E+00 P35679	6.9E+00 P44834	6.9E+00 P34226	6.8E+00 W03412.	6.8E+00	6.8E+00	8.8E+00	6.6E+00	6.6E+00 E	8.6E+00	6.6E+00 C	6.6E+00	6.5E+00 P	6.5E+00 B	6.2E+00	6.2E+00	6.0E+00 B	6.0E+00 A	6.0E+00 A	6.0E+00 A	5.9E+00	5.8E+00 7661	5.7E+00 A	
	Expression	3.58	3.58	0.71	1.28	1.28	8.63	3.28	3.37	1.61	1.92	1.38	0.47	1.53	1.53	82.	3.24	0.72	0.61	2.36	2.36	1.97	7	0.49	1.55	0.5	1.46	0.46	0.67	0.67	7.32	0.99	0.95	
	ORF SEQ ID NO:		28391	1					35729		34011	36107	38125	33623	33624		35969		32216	35827	35828		34931	36067	35488	36337	32717	35565	36274	36275	32193		32816	
L	SEQ ID NO:		15743		19740		- 1	24047	ı		20875	22897	22915	20503	20603	21721	22757	18010	19218	22622	22622	23743	21768	22851	22294	2310g	19871	22367	23057	23057	19196	16270	19752	
	Probe SEQ ID NO:	2977	2977	6931	7049	7049	9498	11359	9882	11215	8184	10249	10267	7808	7808	9031	10109	5202	6450	9974	9974	11073	9079	10203	9842	10460	6936	9716	10411	1041	6428	3514	7061	

Page 5 of 536 Table 4 Single Exon Probes Expressed In Brain

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Mus musculus immunoglobulin scavenger receptor IgSR mRNA, complete cds	DNA POLYMERASE III, ALPHA CHAIN POLC-TYPE (POLIII)	LYCOPENE BETA CYCLASE	HEAT-INDUCIBLE TRANSCRIPTION REPRESSOR HRCA	Mus musculus DNA methyltransferase (Dnmt1) gene, exons 30, 31, and 32	PNEUMOLYSIN (THIOL-ACTIVATED CYTOLYSIN)	Chicken alpha-cardiac actin gene	Chicken alpha-cardiac actin gene	VITELLOGENIN PRECURSOR (VTG) [CONTAINS: LIPOVITELLIN LV-1N; LIPOVITELLIN LV-1C; LIPOVITELLIN LV-21	REP1 PROTEIN	REP1 PROTEIN	RHODOPSIN	RHODOPSIN	Bovine immunodeficiency-like virus surface envelope gene, 5' end of cds	DNA POLYMERASE GAMMA (MITOCHONDRIAL DNA POLYMERASE CATALYTIC SUBUNIT)	Homo sapiens HERPUD1 gene for stress protein Herp, complete cds	PROBABLE ANTIBACTERIAL PEPTIDE POLYPROTEIN PRECURSOR	QV4-HT0691-270400-186-f09 HT0691 Homo sapiens cDNA	Drosophila orlentacea R1B retrotransposable element reverse transcriptase gene, parttal cds	HYPOTHETICAL 61.1 KD PROTEIN C23E2.03C IN CHROMOSOME I	RHODOPSIN	COLICIN N IMMUNITY PROTEIN (MICROCIN N IMMUNITY PROTEIN)	601894910F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4124114 5	601890420F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4131509 5	Canis familiaris skeletal muscle chloride channel CIC-1 (CLCN1) mRNA, complete cds	Mycobacterium tuberculosis H37Rv complete genome; segment 103/162	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis	(HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	Eunice australis histone H3 (H3) gene, partial cds	RC3-CN0042-100800-011-c10 GN0042 Homo sapiens cDNA	PM0-BT0547-310100-002-604 BT0547 Homo sapiens cDNA	601875654F1 NIH_MGC_55 Homo saplens cDNA clone IMAGE:4099716 5'
де схоп Рто	Top Hit Database Source	LN LN	SWISSPROT	SWISSPROT	SWISSPROT	NT	SWISSPROT	TN	NT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	TN	SWISSPROT	IN	SWISSPROT	EST_HUMAN	NT	SWISSPROT	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	ΙN	TN		L	۲	EST HUMAN	EST_HUMAN	EST_HUMAN
	Top Hit Acesslan No.	00 AF302046.1	DO P75080	5.6E+00 Q55278	P47447	5.5E+00 AF175425.1	5.5E+00 P11990	X02212.1	5.4E+00 X02212.1	00 091062	P40379	P40379	Q17094	Q17094	5.3E+00 L43126.1	5.3E+00 P54098	AB034990.1	QZ7905	5.2E+00 BE184840.1	5.2E+00 AF248070.1	Q10136	016005	P09182	DO BF310443.1	5.0E+00 BF308561.1	AF162445.2	5.0E+00 Z83860.1		4.9E+00 U91328.1	4.8E+00 AF185255.1	4.8E+00 BF367909.1	4.8E+00 AW750067.1	4.7E+00 BF240552.1
	Most Similar (Top) Hit BLAST E Value	5.7E+00	5.6E+00	5.6E+00	5.5E+00 P47447	5.5E+00	5.5E+00	5.4E+00	5.4E+00	5 4F+00	5.4E+00 P40379	5.4E+00 P40379	5.4E+00 Q17094	5.4E+00 Q17094	5.3E+00	5.3E+00	5.3E+00	5.3E+00 Q27905	5.2E+00	5.2E+00	5.2E+00 Q10136	5.1E+00 016005	5.1E+00	5.0E+00	5.0E+00	5.0E+00	5.0E+00		4.9E+00	4.8€+00	4.8E+00	4.8E+00	4.7E+00
	Expression Signal	0.95	1.13	2.59	0.69	1.28	3.09	1.14	1.14	1.54	0.83	0.83	1.83	1.83	1.32	3.23	0.49	3.2	0.91	0.95	. 2	6.0	1.19	0.72	0.59	3.07	8.95		0.71	10.86	0.47	5.28	1.86
	ORF SEQ ID NO:	32817			31901				32515		34534			35785	30102			37548						31944		36197	37214				33879		25731
	Exon SEQ ID NO:	19752	20142	23223	18934	23369	23221	19492	19492	20465	21390	21390	22584	22584	17466	20673	21573	24225		22919		21552	22376	18969	22742	22977	23922		22780	16784	20748	21131	13090
	Probe SEQ ID NO:	7061	7468	11456	6157	10678	11454	0830	0830	7769	8698	8698	9636	9836	4734	7978	8882	11628	5377	10271	11150	8861	9725	6193	10094	10330	11260		10132	4039	8054	8439	283

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			_	10						_		_																		
ongje Exon Proces Expressed in Brain	Top Hit Descriptor	801875654F1 NIH MGC 55 Home contract DMA -1	Homo sapiens chromosome 21 sagmant HS21 Coso	7e89g10.xt NCI_CGAP_CLL1 Homosephers CDN done IMAGE:3292098 3' similar to TR:O75140 O75140 KIAA0645 PROTEIN :contains alement PTDs : contains a	7688g10.x1 NCL_CGAP_CLL1 Home saplens Capitality as a contained to TR.O75140 O75140 KIAA0845 PROTEIN	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	Streethought on Doctors	Griedickýsus sp. r Cudous complete genome, 18/27, 2267260-2392728 Archaevalchie fulcidus confine confi	802123238F1 NIH MGC 58 Home contains 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	802077565FF1 NCL CRAP BE ST LICENS CON FINA CHILD STATE CHILD STATE CHILD S	602072686F1 NCI CGAP Bridt Home smiles CDNA close IMA GE:4216284 6	Murine James for MHC class III/ID) concluded in the Constitution of the Constitution o	domo sanjens ne itronhii collacenco (A) CNA)	Plasmodium falciparum R208+uard sens, even 4	Tebonema nallidim section 38 of 01 golds, con-	Homo seplent gutathione S-transferase thete 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	MICROSOMAL DIPEPTIDASE PRECURSOR (MDP) (DEHYDROPEPTIDASE-I) (RENAL DIPEPTIDASE)	AE-A DROTEIN IN VANDLOID WILLIAM PROCESSION	EXTENSIA PRECIDENCE MALL MALL MALL MALL MALL MALL MALL MAL	EXTENSIN PRECIDEND (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN)	W(6/2003 x1 Shares NEI T GEO S4 HOLL TO THE SAME THE SAME SHARES NEI T GEO S4 HOLL TO THE SAME SAME THE SAME SAME SAME SAME SAME SAME SAME SAM	NUBBIN PROTEIN /TWAIN PROTEIN /POLITION AND INTERMACE. 2360692 3	HEXOSE TRANSPORTER HXTR	CELLUI AR TIMOR ANTICENDES	CELLULAR THMOR ANTIGEN BES	801110727F1 NIH MGC 18 Homs senions above 12.	601859030F1 NIH MGC :88 Home Saniens CDNA clone INACE: 3331534 5	YY1 PROTEIN PRECURSOR	Pathopecter vessoensis mRNA for calcina into A complete and	GENE 68 PROTEIN
	Top Hit Database Source	EST HUMAN	N	EST HUMAN	EST HUMAN	į į	F	_LN	EST HUMAN	EST HUMAN	ST HUMAN	Ν	N	N	LN	LN TN	TOGGOSI	T	Т	1	7	Т		SWISSPROT	T	Т	Т	SWISSPROT		/ISSPROT
5	Top Hit Acession No.	-00 BF240552.1	+00 AL163280.2	-00 BE646437.1	-00 BE646437.1		+00 D63999 1		Γ	Γ		00 X13414.1	4.3E+00 AF059679.1		00 AE001222.1	4.3E+00 AF240786.1										-				
	Most Similar (Top) Hit BLAST E Value	4.7E+00	4.7E+00	4.6E+00	4.6E+00	4.6E+00	4.6E+00	4.5E+00	4.5E+00	4.4E+00	4.4E+00	4.4E+00	4.3E+00 /	4.3E+00 Y13402.1	4.3E+00	4.3E+00	4.2E+00 F	4.2E+00 P51828	4.2E+00 P13983	4.2E+00 P13983	4.2E+00 AI809013.1	4.2E+00 P31368	4.2E+00 P40886	4.1E+00 O	4.1E+00 009185	4.1E+00 BE253688.1	4.1E+00 BF247939.1	4.1E+00 O	4.1E+00 AB041523.1	4.1E+00 P28964
	Expression Signaí	1.89	2.38	1.18	1.18	0.61	2.31	2.59	1.78	96.0	96:0	1.66	0.68	2.03	0.65	7.64	3.44	0.87	2.62	2.62	4.68	1.06	0.48	0.56	0.56	0.84	0.65	8.73	0.62	4.32
	ORF SEQ ID NO:		28679	34948	34949			37528	37685	28447	28448			33097	33280	36696		31206	32627	32628	34697	35672		31569	31570	32760	32863	33332	+	33459
	Exon SEQ ID NO:		18030	21783	21783	22935	23724	24204	- 1	- 1	ı	1888	18807	20 19	20186	23453	18229	18305	19591	19591	21550	22469	22697	25079	25079	19704	19799	20229	20345	20347
	Probe SEQ ID NO:	284	3268	9095	9095	10287	11054	11605	11762	3035	3035	9109	6027	7338	7615	10769	5430	5507	6874	6674	8829	9818	10049	5846	2846	7912	1	7559	7681	7883

Page 7 of 536 Table 4 Single Exon Probes Expressed in Brain

Γ	·	٦	7	7	٦		7	\neg	Т	7	П		٦	٦	\neg	7	7		П					П	П	T		Γ
	Top Hit Descriptor	GENE 68 PROTEIN	Pan troglodytes novel repetitive solo LTR element in the RNU2 locus	50S RIBOSOMAL PROTEIN L4	602247938F1 NIH_MGC_62 Homo septens cDNA clone IMAGE:4333209 5'	CYCLIN-DEPENDENT KINASE INHIBITOR 18 (CYCLIN-DEPENDENT KINASE INHIBITOR P27)	(P <i>ZT</i> KIP1)	HYPOTHETICAL PROTEIN HVLF1	601507510F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909051 5'	GLC7-INTERACTING PROTEIN 1	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE ; ISOMALTASE]	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE ; ISOMALTASE]	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE ; ISOMALTASE]	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE ; ISOMALTASE]	CELL DIVISION PROTEIN FTSY HOMOLOG	HYPOTHETICAL PROTEIN KIAA0144	NITRIC-OXIDE SYNTHASE (NOS, TYPE I) (NEURONAL NOS) (N-NOS) (NNOS)	Ureaplasma urealyticum section 33 of 59 of the complete genome	CYTOCHROME C OXIDASE POLYPEPTIDE III	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE GLYCOPROTEIN MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)]	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN	(ENVELOPE GLYCOPROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSI RUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)]	N.tabacum chitinase gene 50 for class I chitinase C	Mus musculus seminal vesicle secretory protein 99 (MSVSP99) gene, promoter region	MR0-BN0070-300500-028-h05 BN0070 Homo sapiens cDNA	MR0-BN0070-300500-028-h05 BN0070 Homo sapiens cDNA	 Human hereditary haemochromatosis region, histone 2A-like protetn gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	I NO STUTTON TO ANISOCIOTIONAL DECLI ATOD IN AIDO DOSE INTERCENIO DECIONI
	Top Hit Database Source	SWISSPROT	NT	SWISSPROT	EST_HUMAN		SWISSPROT	SWISSPROT	EST_HUMAN	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	LN	SWISSPROT	SWISSPROT		SWISSPROT	LZ	TN	EST_HUMAN	EST_HUMAN	Ę	100000000
	Top Hit Acession No.	P28964	U57503.1	P11253	BF692425.1		P46414	P09716	BE885880.1	P38229	062653	062653	062653	062653	033010	Q14157	061309	AE002132.1	P14546	P07564		P07564	X64518.1	AF055466.1	BE814357.1	BE814357.1	U91328.1	000000
	Most Similar (Top) Hit. BLAST E Value	4.1E+00	4.1E+00	4.1E+00	4.1E+00			4.1E+00		4.0E+00	4.0E+00		4.0E+00	4.0E+00	4.0E+00	4.0E+00	4.0E+00	4.0E+00	4.0E+00	4.0E+00		4.0E+00		3.9E+00		3.9E+00	3.9E+00	00.50
	Expression Signal	4.32	2.53	0.57	2.48		0.48	3.06	11.69	0.95	7.0	0.77	0.75	0.75	1.44	0.45	0.4	0.63	1.53	2.27		2.27	4.61	8.24	2.91	2.91	0.55	3
	ORF SEQ ID NO:	33460	33638	35295								32525	32524	32525	32843			35931		37444		37445			31279	31280	32367	l
	SEQ ID NO:	20347	20512	22118	22224	1	22853	23483			19500	19500	19500	19500	19778	L	22494	<u>_</u>		24137		24137	L		1	18369	19354	1
	Probe SEQ ID NO:	7683	7817	9440	9571		10205	10800	10892	3533	5372	6372	6838	6838	7089	8772	9843	10065	11463	11637		11537	3494	4287	5572	5572	6591	67774

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Table 4
Single Exon Probes Exonescad in E

Single Exon Probes Expressed in Brain	lit sise Top Hit Descriptor	Human MHC class II Namboovie aptition (DBurd-bets-1) come 3	X, lasvis mRNA for M4 muscarinic reventhr	Homo sapiens NF2 gene	Helicobacter pylori, strain J99 section 123 of 132 of the complete canonia	Г	T	Г	П	Arabidopsis thaliana DNA chromosome 4. contin fragment No. 30	Thermoplasma acidophilum complete genome: secoment 3/5	Homo saplens glucokinase (haxokinase 4, maturity onset diabetes of the young 2) (GCK), nuclear gene	encoding mitochondrial protein, mRNA	П				Arabidopsis thaliana DNA chromosome 4. contta fragment No. 2	Г	Г	Г	Г	Pseudomonas aaruginosa PA01, section 8 of 529 of the complete genome	Escherichia coli glycerophosphate dehydrogenase (glpD) gene, partial cds; and the translation start site has	been verired (gip⊏), me translation start site has been verified (gipG), and repressor protein (gipR) genes, complete ods	Crotospoddium fells heaf shock protein 20 (HSD20) and a shock	Borella burgdorfert (strein 25015) aufter surface problem of partial conditions	Т	Ι.			\Box	
igle Exon	Top Hit Database Source	Į	ļ.	١	Σ	SWISSPROT	EST HUMAN	EST_HUMAN	LN L	Ę	¥		Į.	FZ	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΙN	NT		Þ	Į	Ę	EST HUMAN	SWISSPROT	EST HUMAN	EST HUMA	NT	
รั ช	Top Hit Acession No.	3.9E+00 M23907.1	3.9E+00 X65865.1	3.9E+00 Y18000.1	AE001562.1	3.8E+00 Q57830	+00 AI493849.1	3.8E+00 D44725.1	3.8E+00 AJ390961.1	+00 AL161539.2	:+00 AL445065.1		4503950 NT	3./E+00 U43541.1	3.7E+00 BF669279.1	3.7E+00 BF669279.1	3.6E+00/AV761055.1	3.6E+00 AL161472.2	3.6E+00 BF316316.1	D12367.1	3.6E+00 D12367.1		+00 AE004447.1		3.6E+00 M96795.1	2			+00 P24557	+00 AA190998.1	+00 AA190998.1	+00 AL161553.2	
	Most Similar (Top) Hit BLAST E Value	3.9E+00	3.9E+00	3.9E+00	3.8E+00	3.8E+00	3.8E+00	3.8E+00	3.8E+00	3.7E+00	3.7E+00		3.7E+00	3./E+00	3.7E+00	3.7E+00	3.6E+00	3.6E+00	3.6E+00	3.6E+00	3.8E+00	3.6€+00	3.6E+00		3.6E+00	3.5E+00	3.5E+00	3.5E+00	3.5E+00	3.5E+00	3.5E+00	3.5E+00	
	Expression Signal	4.3	1.86	3.3	6.0	96.0	0.66	1.1	0.62	12.29	1.03		0.55	0.7	2.23	2.23	5.19	1.06	0.74	0.95	0.95	3.83	3.83		4.07	1.1	1.17	1.18	0.56	1.02	1.02	0.98	
	ORF SEQ ID NO:	33022	34046	36403		32054	32828	34161		29379		_	0.000	34920	3/363	3/364	25986		30498	34280	34281	34378	34379	<u> </u>		28852		31864		34763	34764	35227	
	SEQ ID NO:	19946	20910						_]		19757	2400	24785	2017	75057	7400	13359	17477	17983	21142	21142	21235	21235		23444	16003	18695	18896	21076	21621	21621	22055	
	SEQ ID NO:	7262	8216	11365	2635	6297	6873	8331	9694	4004	7068		8008	200	1408	9	579	4745	5174	8420	8450	8543	8643		10759	3241	5911	6118	888	8930	8930	9393	

Page 9 of 536 Table 4 Single Exon Probes Expressed in Brain

Γ			٦							7	丁	T	٦	7	٦	7	٦	<u>-</u>	П		Ţ	T	-	T	7	٦	7	٦	٦	٦	٦	
	Top Hit Descriptor	Bos taurus mRNA for Ran-binding protein 2, partial	Brassica napus RPB5d mRNA, complete cds	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT	Human atternatively spliced potassium channels ROM-K1, ROM-K2, ROM-K3, ROM-K4, ROM-K5, and ROM-K6 (KCNJ1) gene, complete cds	Homo saplens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 2/3	Homo sapiens partial TM4SF2 gene for tetraspanin protein, exon 6	Saccharomyces cerevisiae MSS1 gene, complete ods	Homo sapiens DiGeorge syndrome critical region, centromeric end	PUTATIVE IRON ALCOHOL DEHYDROGENASE	PUTATIVE IRON ALCOHOL DEHYDROGENASE	Homo sapiens serine palmitoy transferase, subunit II gene, complete cds; and unknown genes	Bacilius halodurans genomic DNA, section 5/14	Becilius halodurans genomic DNA, section 5/14	D.renio zp-50 POU gene	D rento zp-50 POU gene	Homo sapiens carcinoembryonic antigen-relatad cell adhesion molecule 1 (billary glycoprotein) (CEACAM1), ImRNA	SQUALENE-HOPENE CYCLASE	SQUALENE-HOPENE CYCLASE	PHOSPHOGLYCERATE KINASE, CYTOSOLIC	PHOSPHOGLYCERATE KINASE, CYTOSOLIC	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4	VON WILLEBRAND FACTOR PRECURSOR (VWF)	Chlamydomonas reinhardtil chloroplast DNA for rps9, ycf4, ycf3, rps18 genes	Chlamydomonas reinhardtii chloroplast DNA for rps9, ycf4, ycf3, rps18 genes	PERIPLASMIC (NIFE) HYDROGENASE SMALL SUBUNIT (NIFE HYDROGENLYASE SMALL CHAIN)	Scerevislae threonine deaminase (ILV1) gene, complete cds	Oryzias latipes OIGO6 gene for guanylyl cyclase C, complete cds	Sus scrofa choline acetyltransferase gene, promoter region	IHYPOTHETICAL 142.5 KD PROTEIN C23E2.02 IN CHROMOSOME I
ואים העבווו	Top Hit Database Source	NT	IN	SWISSPROT	SWISSPROT	FX	N	NT	NT	IN	SWISSPROT	SWISSPROT	NT	NT	님	NT	Ŋ	ΙN	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	NT	NT	SWISSPROT	NT	NT	NT	TOGGSSIWS
5	Top Hit Acession No.	3.5E+00 AJ133723.1	3.4E+00 AF254577.1	P04052	P04052	3.4E+00 U65406.1	00 AJ229042.1	00 AJ250587.1	3.4E+00 AF013167.1	00 L77570.1	00 Q09669	00 Q09669	00 AF111168.2	00 AP001511.1	00 AP001511.1	X96422.1	X96422.1	4502404 NT	00 P54924	P54924	00 P12783	00 P12783	P18931	P18931	P04275	Y13655.1	Y13655.1	00 P13061	00 M36383.1	00 AB016081.2	L33836.1	040438
	Most Similar (Top) Hit BLAST E Value	3.5E+00	3.4E+00	3.4E+00	3.4E+00 P04052	3.4E+00	3.4E+00	3.4E+00	3.4E+00	3:4E+00	3.3E+00	3.3E+00	3.3E+00	3.3E+00	3.3E+00	3.2E+00 X96422.1	3.2E+00	3.2E+00	3.2E+00	3.2E+00 P54924	3.2E+00	3.2E+00 P12783	3.2E+00		١			3.2E+	3.2E+00	3.2E+00	3.2€+00 \∟33836.1	2 4 1 100 04042
	Expression Signal	0.46	2.94	2.64	69.0	0.7	0.67	0.54	2.97	1.89	1.57	1.57	0.79	6.0	6.0	1,64	6.0	1 08	1.06	1.08	2.7	2.7	1.78	1.78	7.0	2.65	2.65	4.51	0.87	2.03	2.44	27.0
	ORF SEQ ID NO:	36283	26933	L	33374		34813			37429						25908					31214	31215					33449					2000
	Exon SEQ ID NO:	23063	14247	19945	20267	21289	21662		22812	24119	18759	18759	20489	23008	23008	13273	l	17413	1	18280	18313	18313	18988			1	l	21619	22108	22689	24500	ł
	Probe SEQ ID NO:	10417	5	7261	7601	8577	8972	8 5	10164	11519	5977	5977	7794	10361	10361	88	4004	4679	288	5481	5515	5515	6214	6214	7505	7872	7672	8928	9430	10041	11946	100

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	Most Similar (Top) Hit Acession (Top) Hit Database Signal BLASTE No. Source	0.94 3.1E+00 AF303225.1 NT Bacillus alcalophilus pectate (vase (belE) gene, complete cds	3.1E+00 P40985 SWISSPROT	3.1E+00 P49894 SWISSPROT	3.1E+00 P49894 SWISSPROT	SWISSPROT	3.1E+00 7524759 NT	3.1E+00 Q10126 SWISSPROT	3.1E+00 P49365 SWISSPROT	SWISSPROT	00 S56860.1	3.0E+00 8923984 NT	3.0E+00 X53096,1 NT	3.0E+00 X56037.1 NT	μN	3.0E+00 P18406 SWISSPROT	O Q13201 SWISSPROT	┱	0 Q58605	1.57 3.0E+00 Q16181 SWISSPROT CDC10 PROTEIN HOMOLOG	RETINAL GUANYLYL CYCLASE 2 PRECURSOR (GUANYLATE CYCLASE 2F, RETINAL) (RETGC-2) (ROD OUTER SEGMENT MEMBRANE GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE 2)	3.0E+00 P51842 SWISSPROT	.2 NT	2.9E+00 Z36879.1 NT	2.9E+00 O14514 SWISSPROT
	Most Sim (Top) H BLAST Value	3.1E+	L					•			3.1E+	3.0E+				L		3.0E+(3.0E+(
-	ORF SEQ ID NO:		20681 33807		21192 34334	21889	447 35652	338	382 36095	20	99			32228		46	.85	97		40 36474	11 36860		40 27464		32861
	Probe Exon SEQ ID SEQ ID NO: NO:	7627 202		l	1			9888 22538		11440 23207	11463 24066	2842 15610					7096 19785	8805 21497	10192 22840	- 1	10931 23611	 10931 23611	2004 14740	ı	7110 19798

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Table 4
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7110	19798	32882	5.21	2.9E+00	014514	SWISSPROT	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 1 PRECURSOR
7356	20037	33115	6.84		P46589	SWISSPROT	ADHERENCE FACTOR (ADHESION AND AGGREGATION MEDIATING SURFACE ANTIGEN)
7787	20463	28565	0.67	2 85+00	P05844	SWISSPROT	STRUCTURAL POLYPROTEIN [CONTAINS: MAJOR STRUCTURAL PROTEIN VP2; NONSTRUCTURAL PROTEIN VP4: MINOR STRUCTURAL PROTEIN VP31
							STRUCTURAL POLYPROTEIN [CONTAINS: MAJOR STRUCTURAL PROTEIN VP2;
7767	20463		0.67		P05844	SWISSPROT	NONSTRUCTURAL PROTEIN VP4; MINOR STRUCTURAL PROTEIN VP3
7996	20691	33819	1.03		BF344171.1	EST_HUMAN	602017413F1 NCI_CGAP_Bm64 Homo saplens cDNA clone IMAGE:4153059 5
1440	14187	26872	4.4	2.8E+00	AF186398.1	IN	Buxus harlandii maturase K (matk') gene, partial cds; chloropiast gene for chloropiast product
1629	14375		2.74		AL181552.2	TN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 52
7207	19892	32968	5.72		8393724 NT	TN	Mus musculus endomucin (LOC63423), mRNA
9513	22168		0.54	2.8E+00	BE565182.1	EST_HUMAN	601342758F1 NIH_MGC_63 Homo sapiens cDNA clone IMAGE:3684807 6
10588	19892	32968	1.32	2.8E+00	8393724 NT	NT	Mus musculus endomucin (LOC63423), mRNA
224	13036	25672	13.51	2.7E+00		IN	Mus musculus per-hexamer repeat gene 3 (Phxr3), mRNA
224	13036	25673	13.51	2.75+00	LN 9056799	LN	Mus musculus per-hexamer repeat gene 3 (Phxr3), mRNA
5 <u>4</u> 84	18263	31154	1.17	2.7E+00	L14005.1	TN	Homo sapiens apoA polymorphism Kringle IV gene, exons 1 and 2
8045	20739		9.0		U15947.1	LN	Ipomoea purpurea chalcone synthase (CHSB) gene Including complete 5'UTR and complete cds
8867	21558		1.83	2.7E+00	AL116459.1	ΙN	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
0833	20403	33510	0.73	00+ <u>34</u> 6	1 POLABOWA	NAMIN TRE	xx88e12.x1 NCI_CGAP_Bm35 Homo sapiens cDNA clone IMAGE:2591374 3' similar to gb:M17733 THYMOSIN BFTA-4 (HUMAN)
10397	23043		1.75	2.7€+00		EST HUMAN	CM0-BT0281-031199-087-h04 BT0281 Homo sapiens cDNA
4626	17361	29994	5.15	2.6E+00	AF068749.1	NT	Mus musculus sphingosine kinase (SPHK1b) mRNA, complete cds
5460	18259	31149	1.68		6755601 NT	F	Mus musculus SRY-box containing gene 13 (Sox13), mRNA
5480	18259	31150	1.68	2.6E+00	1N +099529	LN T	Mus musculus SRY-box containing gene 13 (Sox13), mRNA
5736	18528		0.59	2.6E+00	Y17062.1	NT	Mycobacterium fortultum furA II gene
7454	25424		0.82	2.6E+00	AJ224639.1	NT	Homo saplens Surf-5 and Surf-6 genes
7600	20266		6.04	2.6E+00	AF235502.1	IN	Mus musculus SH2-containing inosital 5-phosphatase (Ship) gene, exons 16 through 27, and complete ods
7958	20853	33778	1.13	2.8E+00	AJ132180.1	NT	faba bean necrotic yellows virus C2-Eg gene, Isolate Egyptian EV1-83
7958	20653	33777	1.13	2.6E+00	AJ132180.1	NT	faba bean necrotic yellows virus C2-Eg gene, isolate Egyptian EV1-83
9557	22210	35395	2.83	2.6E+00	AL161540.2	IN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 40
10253	22901		1.67		9055193 NT	NT	Mus musculus cleavage and polyadenylation specificity factor 3 (Cpsf3), mRNA
10953	23630	36878	1.32	2.6E+00	AF143675.1	NT	Hantavirus Z10 segment M G1/G2 glycoprotein (Z10) gene, complete cds
12560	25304		3.17	2.6E+00	11419220 NT	NT	Homo saplens A TP-binding cassette, sub-family B (MDR/TAP), member 4 (ABCB4), mRNA
1448	14195	28878	3.73	2.5E+00	AJ271844.1	N-	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4

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		T	Т	T	Т	Г	Т	Т	Т	Т	Т	Т	Т	Т	Т	1	Т	Г	Г	т	_	1	Ι-	г	_	Т	Т	_	Τ-	1		Т	$\overline{}$
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4	TEICHOIC ACID BIOSYNTHESIS PROTEIN F	TEICHOIC ACID BIOSYNTHESIS PROTEIN F	TEICHOIC ACID BIOSYNTHESIS PROTEIN F	TEICHOIC ACID BIOSYNTHESIS PROTEIN F	Vibrio cholerae ctxA gene and ctxB gene for cholera toxins, complete cds	QV4-FT0005-110500-205-907 FT0005 Homo saplens cDNA	Homo sepiens clethrin, heavy polypeptide-like 1 (CLTCL1) mRNA	Rice DNA for aldolase C-1, complete cds	601175779F1 NIH_MGC_17 Hamo sapiens cDNA clone IMAGE:3531090 5'	DNAJ PROTEIN	Mus musculus EIF4H gene, partial cds; LIMK1 gene, complete cds; and ELN gene, partial cds	Chicken alpha-3 collagen type VI mRNA, 3' end	Homo sapiens double C2-like domains, alpha (DOC2A) mRNA	VITELLOGENIN I PRECURSOR (YOLK PROTEIN 1)	602120856F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4278012 5'	602120856F1 NIH_MGC_56 Homo saplens cDNA done IMAGE:4278012 5'	CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)	OD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)	Helicobacter pylori, strain J99 section 47 of 132 of the complete genome	RC2-PT0004-031299-011-d05 PT0004 Home saplens cDNA	ENDOCHITINASE B PRECURSOR (CHN-B)	SKIN GRANULE PROTEIN PRECURSOR	SKIN GRANULE PROTEIN PRECURSOR	H.sapiens CTGF gene and promoter region	XMULOSE KINASE (XYLULOKINASE)	hre3706.x1 NCI_CGAP_Kid11 Hamo sepiens oDNA clane IMAGE:3133187 3'	hr63f06.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3133187 3'	DENITRIFICATION REGULATORY PROTEIN NIRQ	Fregaria x enanassa cytosolic ascorbate peroxidase (ApxSC) gene. ApxSC-o alleie, complete cds	G.domesticus artificial single chain antibody gene (L3)	Bos faurus partial cytb gene for cytochrome b
gle Exon Prol	Top Hit Detabase Source	ΤN	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	LN	EST_HUMAN	LN	IN	EST_HUMAN	SWISSPROT	NT	LN	NT	SWISSPROT	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	NT	EST_HUMAN	SWISSPROT	SWISSPROT	SWISSPROT	LN	SWISSPROT	EST_HUMAN	EST_HUMAN	SWISSPROT	LΝ	NT	LV
Sin	Top Hit Acession No.	-00 AJZ71844.1	213485	-00 P13485	-00 P13485	-00 P13485	H00 D30052.1	-00 AW949158.1	4502902 NT	-00 D50307.1	2.5E+00 BE297758.1	240170	2.5E+00 AF289665.1	M24282.1	4503352 NT	2.4E+00 P02843	3F667502.1	2.4E+00 BF667502.1	26842	26842	2.4E+00 AE001486.1	2.4E+00 AW876128.1	24091	13673	-00 P13673	(92511.1	00 P09099	00 BE326702.1	2.4E+00 BE326702.1	00 Q51481	2.4E+00 AF158662.2	746724.1	2.3E+00 AJ401081.1
	Most Similar (Top) Hit BLAST E Value	2.5E+00	2.5E+00 P13485	2.5E+00	2.6E+00	2.5E+00	2.5E+00	2.5E+00/	2.5E+00	2.5E+00	2.5E+00	2.5E+00 P40170	2.5E+00	2.4E+00 M24282.1	2.4E+00	2.4E+00 F	2.4E+00	2.4E+00	2,4E+00 P26842	2.4E+00 P26842	2.4E+00	2.4E+00 /	2.4E+00 P24091	2.4E+00 F	2.4E+00 F	2.4E+00 X92511.1	2.4E+00 F	2.4E+00 E	2.4E+00 E	2.4E+00 C	2.4E+00/	2.3E+00 Z46724.1	2,3E+00 A
	Expression Signal	3.73	2.22	2.22	1.63	1.63	0.64	0.99	0.58	1.53	0.67	1.34	3.08	1.13	6.09	4.16	0.78	0.78	2.4	2.4	2.63	1.61	7.36	2.56	2.56	1.86	6.55	1.62	1.62	0.87	2.16	13.6	1.35
	ORF SEQ ID NO:	26879	31434	31435	31434	31435	32406	33431	33477	34841	35608			28428	30203	31657	33040	33041	33865	33866			34563	35788	36789	35868		38079	36080	36364	37327	26650	
.	Exon SEQ ID NO:	14195	18616	18515	18515	18515	19392	20323	20383	21691	22403	24128	24498	15778	17579	18705	19964	19964	20734	20734	20804	21241	21419	22586	22588	22655	22791	22868	22868	23136	24022	13980	16845
	Probe SEQ ID NO:	1448	6723	6723	6367	6367	6830	7659	7700	9001	9762	11528	11943	3012	4849	5920	7280	7280	8039	8039	8110	8549	8727	9938	9938	10007	10143	10220	10220	10490	11331	1231	4102

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					Builo	10 EXUIT FLUX	Single Exort Probes Expressed in the community of the com
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
	<u> </u>			100		FST HUMAN	J7340F Human fetal heart, Lambda ZAP Express Homo sapiens conv. Gordon J7340 Samina C. PROLYLCARBOXYPEPTIDASE
5744	l			2.35+00 1402+5.1	5978554		Rattus norvegicus ATPase, Ca++ transporting, ubiquitous (Atp2a3), mRNA
7354		33113		2.35-100		/ISSPROT	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENT-D)
7495	25425			Z.3E+00 P07 199		±14	M mazei dnak and dnaJ genes homologues coding for Dnak and DnaJ
7879	20343		1.01	2.3E+00			Polymens omaticings mitochondrion, complete genome
8006	ı	34849		2.3E+00	1N / LEGESS		A DIA 44 STEIN ICOSYLTRANSFERASE (GALACTOSIDE 3-L-FUCOSYLTRANSFERASE)
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Topics Explanation III Drain	Top Hit Descriptor	bb17h12.x1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:2963207 3' similar to ch-Dusase Manages	mRNA for nuclear pare-targeting-complex component of (MOUSE);	001094/33F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3948661 5	manustruston 171 PROTEIN A qm89b03.x1 Soares_placenta_8te9weeks_2NbHP8to9W Homo saniens cDNA_class NAA OE Assasses	similar to gb: Y00433 GLUTATHIONE PEROXIDASE (HUMAN); qm89b03.x1 Soares, placenta 8toSweeks, 2NbHP8hodW Home seeles, CANA	similar to gb: Y00433 GLUTATHIONE PEROXIDASE (HUMAN);	601855591F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE 4075391 #	Homo sapiens ovarian granulosa cell 13.0 kDa protein hGR74 homolog mRNA complete calc	UROMODULIN PRECURSOR (TAMM-HORSFALL URINARY GLYCOPROTEIN) (THP)	EARLY E1A 28 KD PROTEIN	Mus musculus pre-T cell receptor alpha gene, enhancer region and upstream region	UI-H-BI3-akt-e-08-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2734660 31	HYPO I HE TICAL PROTEIN MG302 HOMOLOG	ALYHA-Z-HS-GL/COPROTEIN PRECURSOR (FETUIN-A)	TRANSCRIPTION INITIATION FACTOR TEILD ALLINAAN.	AU123830 NT2RM2 Home conference ONE	Homo sanions a Capacidal Molymers Curve Clane N ZKMZ000671 6	rome captures por part of the	Orredom 22 Date (UOKDEL) mRNA, complete cds	OF TATILIE BONA AFFILM TO THE BOOM STATE OF THE BONA BOTT TIME BONA AFFILM TO THE BONA AF	R DOWNSTILL MANGET IN THE TANKE OF THE CONTROL OF T	R.norvegicus mRNA for collagen sinhed than I	h13505x1 NCI_CGAP_GUI Anno saplens GNA clone IMAGE 2972168 3' similar to gb:X01877	hi13co527 NOI_COAP GUT Home saplens oDNA clone IMAGE; LIVER (HUMAN);	STRUCTURAL POLYPROTEIN [CONTAINS: NUCLEOCAPSID PROTEIN C. MEMBEANE	GLYCOPROTEINS E1 AND E2]	Escherichia coli 0157 DNA, map position at 46 min., complete cds	Escherichia coli 0157 DNA, map position at 46 min., complete cds	escherchia coli 0157 DNA, map position at 46 mln., complete cds
101 1 10V= 0.6.	Top Hit Database Source		EST HUMAN	Т	T	\top	\neg	HOMAN	٦.	Τ.	SWISSEROI		7	SWISSPRO!	\top		T HUMAN	Т			/ISSPROT	Т	N.	EST HUMAN G			ISSPRO	- LE		
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Columbia Columbia		Expression Signal	0.75	3.2	0.63	, r	9.6	,	0000	256	0.81	1.77	1.77	3.28	1.11	1.11	0.91	1.13	0.59	9.0	2.3	0.59	0.69	2.18	2.18	1.16	0.48	2.57	9		1.84	18.61
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Olingie Exon Probes Expressed in Drain	Top Hit Database Source	Homo saplens small proline-rich protein (SPRR3) gene, exons 1, 2, and 3 and complete cds	Mus musculus ST6Gal/Acill gene, exon 2	B.napus gene encoding endo-polygalacturonase	zd25f01.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:341689 5' similar to EST_HUMAN gb:D29805 N-ACETYLLACTOSAMINE SYNTHASE (HUMAN);	EST_HUMAN 60218609511 NIH_MGC_45 Homo eaplens cDNA clone IMAGE:4310591 3'	Homo sapiens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds	Homo sapiens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds	Uroteuthis chinensis cytochrome c oxidase subunit I (COI) gene, mitochondrial gene encoding mitochondrial protein, partial cds	Uroteuthis chinensis cytochrome c oxidase subunit I (COI) gene, mitochondrial gene encoding mitochondrial	probain, partial cds	Mus musculus ST6GalNAcill gene, exon 2	Mus musculus ST6GalNAcIII gene, exch 2	Brachydanio rerio MHC class II DA-beta-2'01 gene, 3' end	Homo sapiens transglutaminase type I (Tgasel) gene, promoter region		EST_HUMAN UI-H-BIZ-ehr-b-04-0-UI.s1 NCI_CGAP_Sub4 Homo septens cDNA clone IMAGE:2727611 3:		SWISSPROT VIRULENCE FACTOR MVIN HOMOLOG	Mus musculus SIL, MAP_17, CYP_a, SCL & CYP_b genes	Homo sapiens hypothetical protein PRO0971 (PRO0971), mRNA		EST_HUMAN 601283925F1 NIH_MGC_44 Homo sapiens CDNA clone IMAGE::3605647 5	M.musculus COL3A1 gene for collagen alpha-l	M.musculus COL3A1 gene for collagen alpha-l		EST_HUMAN ph6b6_18/1TV Outward Alts-primed hncDNA library Homo sapiens cDNA clone ph6b6_18/1TV	Drosophila melanogaster signal transducting adaptor protein (STAM), serine threonine kinase Ial (IAL), and	П		EST_HUMAN QV4-LT0016-090200-100-d07 LT0016 Homo sepiens cDNA	\neg	Glugea plecoglossi beta-tubulin 2 (btub2) gene, partial cds
eignic	Top Hit Acession No.	10 AF077374.1 NT		18373.1 NT			155827.1 NT	155827.1 NT	-075394.1 NT		1 .	11344.1 NT		H808.1	-005631.1 NT					1297131.1 NT	11437222 NT	37222			52046.1 NT								F162084.1 NT
	Most Similar (Top) Hit BLAST E Value	1.6E+00 Af	1.6E+00 Y11344.1	1.6E+00 X98373.1	1.6E+00 W58426.1	1.6E+00 BF570077.1	1.6E+00 AF155827.1	1.6E+00 AF155827.1	1.6E+00 AF075394.1		1.6E+00 AF075394.1	1.6E+00 Y11344.1	1.6E+00 Y11344.1		1.6E+00 AF005631.1	1.6E+00 BF380703.1	1.6E+00 AW294881.1	1.6E+00 BE697267.1	1.6E+00 Q46378	1.6E+00 AJ297131.1	1.6E+00	1.6E+00	1.6E+00 BE388331.1	1.6E+00 X52046.1	1.6E+00 X52048.1	1.6E+00 A	1.6E+00 T41290.1			1.6E+00 A		1.6E+00 AF037352.1	
	Expression Signal	3.75	1.54	1.24	1.61	5.66	1.9	1.9	0.84		0.84	2.86	2.86	2.16	0.79	0.69	1.06	2.73	1.19	3.28	0.83	0.83		1.94	1.94	0.58	1.32		0.5	1.15	1.15	0.47	0.45
	ORF SEQ ID NO:		27505		28377		29682		30277					L	31543	32146	32387	32901		34112	34636	34637			33550		35480		35911	35947			36365
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	Probe SEQ ID NO:	2038	2042	2282	2961	4011	4319	4319	4842		4942	5024	5024	5737	5823	8378	6810	7145	7929	8277	8428	8428	8970	0986	9360	9487	9834		10047	10085	10085	10242	10491

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	CAPSID PROTEIN PAGICON TANKS. AGGINATION 775	nc16b02.s1 NCI_CGAP_PrI Homo septens cDNA clone IMAGE:1008287 similar to contains.	repetitive element;	Homo sapiens transglutaminase type I (Tgasel) gene, promoter region	Homo sapiens unknown mRNA	Rattus norvegicus jun dimerization protein 2 (jdp-2) mRNA, complete cds	Untarrydophila pneumoniae AR39, section 32 of 94 of the complete genome	Mus musculus a disintegrin and metalloproteinase domain (ADAM) 15 (metaraidin) (Adam18) mBNA	Potato virus A RNA complete genome, Isolate U	Mus musculus I -cell lymphoma invasion and metastasis 1 (Tlam1), mRNA	Potato virus A RNA complete genome, Isolate U	Delnooccous radiodurans R1 section 82 of 229 of the complete chromosome 1	tf2f10.x1 NCI_CGAP_GC6 Hamo saplens cDNA clone IMAGE:2240587 3' similar to TR:O00237 O00237 HKF-1	#12f10.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2240587 3' similar to TB-Onora Consess	HKF-1.;	9910e02.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:31693 6	ON 1476743FT NIH MGC 68 Homo saplens cDNA clone IMA GE:3881555 5	HTPOLHE IICAL 118.4 KD PROTEIN IN BAT2-DALS INTERGENIC REGION PRECLIRSOR	AT PUT HE LICAL 118.4 KD PROTEIN IN BATZ-DALS INTERGENIC REGION PRECURSOR	and 10.51 Soates, tests_NHT Homo sapiens cDNA clone IMAGE:1407115 3/ and 7611.51 Stratagene schizo brain S11 Homo sapiens cDNA	9b:S95936 SEROTRANSFERRIN PRECURSOR (HUMAN):	Homo sapiens WDR4 gene for WD repeat protein, complete cds	DVI 309386F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911181 6	Mouse germine (gM chain gene, mu-delta region	Fromo sapiens hGPIb alpha gene for platelet glycoprotein ib alpha, complete cds	COLOSZOSZTI NIH_MGC_67 Homo saplens cDNA clone IMAGE:4096136 6'	Mushori, r. Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:147697 6	4V3-C1018Z-261099-008-d09 CT0192 Homo saplens cDNA	RCU-1N00/8-150800-034-g05 TN0078 Homo seplens cDNA	2638q06.11 Soares retina NOALUB H.	Section of the sectio
gle Exon Pro	Top Hit Database Source	SWISSPROT	FOU	ES - HOMAN	IN IN	114	I I					114	Z	EST_HUMAN	ı	- 1	FOT THEMAN	Т	7	T	T	T-HUMAN	Т	NAMOL			Т	Т	7	T	Т	7
Sin	Top Hit Acession No.	P54817	F+00 A4246387 4	=+00 AF005634 4		I	,	8	07 0250 I	1.5E+00 A878350 NOT	3	T	T	+00 AI685301.1	+00 Alessan1 1	T	-	T		1 05		+00 Al003254.1 E	T	T	T	T	T	1	T	T		
	Most Similar (Top) Hit BLAST E Value	1.6E+00 P54817	1.6F+00	1.6E+00	1.6E+00	1.5E+00	1.5E+00	1.5F+00	1.5F+00	1.5E+00	1 5F+00 /	1.5F+00/	201	1.5E+00/	1 5F+00 A	1.5E+00 F	1.5E+00 B	1.5E+00 P	1 5F+00 P	1.5E+00 A		1.0E+00 A	1.5E+00 RF887448 4	1.5E+00 K02438 4	1.5E+00 AB038518 4	1.5E+00 BF217818 1	1.5E+00 R81928 1	1.5E+00 AV	1.5E+00 BF376754 1	1.5E+00 BF337944.1	1.5E+00 AA	
	Expression Signal	1.95	1.27	5.27	3.46	5.31	22	2.03	1.95	2	1.85	0.72		0.83	0.83	3.02	1.37	23.98	23.98	0.61	97.0	0.70	0.89	0.84	0.48	0.46	0.62	1.39	6:38	1.77	1.66	
	ORF SEQ ID NO:	36602	36657	31543		25476	25674		27867	27975	27867	28785		31350	31351	32068		32814	32815	33006	33257		33848	34377		34876	35217	35374	35618		35958	
	SEO ID NO:	23361		Li	24300	12859	13037	13384	15131	15235	15131	16127		18437	18437	19083	19720	19751	19751	19930	20165	20390	20716	21234	21605	21722	22045	22188	22411	22600	22744	
	Probe SEQ ID NO:	10670	10728	10747	11705	31	225	909	2410	2519	3135	3368		5642	5642	6312	7028	2060	7060	7245	7463	7727	8021	8542	8914	9032	_[9952	10098	
					-									1_				-		_1		L		_1	_1	_	_1	L			_	

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Single Exon Probes Expressed in Brain	То́р Hit Descriptor	802223425E4 NIII 1100 0011	IN ELITAGE CONTROL STATE HOME SEPTENS CON CIONE IMAGE: 4288137 6'	III. E LITATOR 201009-008-CO4 HT0198 Homo sapiens cDNA	Pondalis Control (1998-004-004-004-004-004-004-004-004-004-00	Pandorina colemaniae chloroplast rbcL gene for ribulose bisphosphate carboxylase, partial cdo	236e09.rt Soares, NNHMPu_S1 Homo sapiens cDNA clone IMAGE:665512 6' similar to contains element	MENAZ repeutive element;	From Saprets Arecel mrnA for AIRE-1, complete cds	601655184R1 NIH MGC 65 Homo saplens cDNA clone IMA 0E:3845805 3'	Printmodustis cartniif on rath quantum muchaelds Lines	The second secon	Preumocystis cannil f. sp. ratti guanine nucleotide binding protein alpha subunit (pcg1) gene, complete cds	M missels trained Live Citionosome 4, config fragment No. 12	Caribarelling on a care of the caribar of the care of	Home soules and the use too rank gene, isolate liber	Home spries of the painting pseudogene for hair keratin, exons 2 to 7	Homo saniene vincienza za central de 7 (14272) (ZNF157) mRNA	Coly lacound a thy distribution and the color of the colo	Chlamydia murdarum, section 66 of 85 of the complete cds	Cyprinus carplo MRPb and MASPb genes for mannose-binding lectin-essociated serine protease (MASP)	AMININAL PHA CHAIN BEED DESCRIPTION	ENTERIOR DE LIGIT DE CITATION PRECURSOR (LAMININ A CHAIN)	Mus musculus alpha-specifin 1 and though (Sanda Texts) — BAYA	Fugu rubripes gamma-aminobutyrio addi receptor bate subunit gene, partial cds; 55kd erythrocyte membrane protein (P55), synaptic vesicle-essociated interest membrane protein (P55), synaptic vesicle-essociated interest membrane protein (YAMD 4)	enhancer protein (PCOLCE) genes, complete c>	PHENOL HYDROXYLASE P3 PROTEIN (PHENOL 2-MONOOXYGENASE P3 COMPONENT)	Human estradiol 17 beta-dehydrogenase gene, complete ods	0VZ 148264F1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4309095 5' PMO-CT0289-291199-004-08 CT0789 Homo sapiens CDNA
gle Exon Pro	Top Hit Database Source	EST HIMAN	ENT LINAN	EST LINES		LZ	HOT HIMAN	1000	EST HIMAAN	EST HUMAN	IN	. <u>‡</u>		Ė	L	FZ	1	L	N	LN	F	SWISSPROT	EST HIMAN	1			ISSPROT	T CI BAAAN	Т
รัฐ	Top Hit Acession No.	+00 BF575545 1	+00 RF145374 4	1	ŀ		AA195528 1	+00 AR006682 1		T			2		_		450799R			1.3E+00 AE002338.2 N		1.3E+00 P25391	T	1.3E+00 6755621 NT		-	ŀ	1	1.3E+00 AW362834.1 E
	Most Similar (Top) Hit BLAST E Value	1.4E+00	1.4E+00	1.4E+00	1.4E+00	1.4년+00	1.4E+00	1.4E+00	1.4E+00	1.4E+00	1.4E+00	1 45+00	1.4E+00	1.3E+00	1.35+00	1.3E+00 Y19213.1	1.3E+00	1.31.+60	1.3E+00	1.3E+00,	1.3E+00/	1.3E+00 F	1.3E+00 E	1.3E+00		1.35 +00	1.3E+00 P19732	13F+00	1.3E+00 A
	Expression Signal	0.79	0.61	0.61	1.06	1.06	1.34	6.16	4.42	4.42	3.46	3.46	1.48	1.81	3.42	20.26	13.71	13.71	1.26	2.27	-	1.27	1.75	0.73		0.89	80: 80:	0.81	7.57
	ORF SEQ ID NO:	35968			36291		36931	37139	37288	37289	37357	37358		-	26320		26692	26693				27862		28354	7000	70000	31330	31590	31667
	SEQ ID NO:	ш		22799	23070	ll	23675	23853	23988	23988	24053	24053	25256		13651	-	14024	14024	14083	14351	14967	15126	15268	15705	6000	18228	18418	18850	18712
	Probe SEQ ID NO:	10108	10151	10151	10424	10424	11003	11188	11381	11381	11404	11404	12079	557	882	1107	1274	1274	1334	- - - - - - - - - - - - - - - - - - -	2239	2405	2553	2940	3584	5427	5622	5863	5928

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Velue	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
828	18712	31668	7.57	1.3E+00	DO AW362834.1	EST_HUMAN	PMo-CT0289-291199-004-f08 CT0289 Homo saplens cDNA
6	19093			1.3E+00	DO M33496.1	F	D.melanogaster no-on-transient A gene product, complete cds
6652	19414					SWISSPROT	HYPOTHETICAL GENE 64 PROTEIN
6739	19573	32606			1.3E+00 M13918.2	LN	Homo saplens fibronectin receptor alpha-subunit precursor (ITGA5) mRNA, partial cds
6854	19554	32584	1.17	1.3E+00	DO BE538819.1	EST_HUMAN	601061420F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3447965 5'
2000	19692	32743	0.81	1.3E+00	00 BE243571.1	EST_HUMAN	TCBAP1D0959 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens cDNA clone TCBAP0959
7358	20039		1.01	1,3€+00		SWISSPROT	ACYLPHOSPHATASE, ORGAN-COMMON TYPE ISOZYMES A AND B (ACYLPHOSPHATE PHOSPHOHYDROLASE)
둖	20891		1.28	1.3E+00	12.1	LN	Sus scrofa plp gene
8346	21039		2.78			EST_HUMAN	601657145R1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866195 3'
8459	21151	34294	0.86		1.3E+00 BE974280.1	EST_HUMAN	601680250R2 NIH_MGC_83 Hamo saplens cDNA clane IMAGE:3950532 3'
8611	21303		1.78	1.3E+00	1N 470168	NT	Homo saplens GL004 protein (GL004), mRNA
8689	21381	34525	0.79	1.3€+00	1.3E+00 Al927629.1	EST_HUMAN	wo85a07.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2462100 3'
9415	22093		5.24	1.3E+00	00 AF042084.1	IN	Homo sapiens heparan glucosaminyl N-deacetylase/N-sulforansferase-2 gene, complete cds
9424	22102	35273	2.56	1.3E+(NT	S.alba phr-1 mRNA for photolyase
24	22102		2.56	1.3E+(00 X72019.1	NT	S.alba phr-1 mRNA for photolyase
24	22177		96.0	1.3E+(00 AF059250.1	LNT	Homo sapiens lipoxygenase (ALOX12B) mRNA, complete cds
9569	22222	35407	1.56	1.3E+00	0 000754	SWISSPROT	LYSOSOMAL ALPHA-MANNOSIDASE PRECURSOR (MANNOSIDASE, ALPHA B) (LYSOSOMAL ACID ALPHA-MANNOSIDASE) (LAMAN)
9651	22303	35498	1.14	1.3E+00	IO A1927629.1	EST_HUMAN_	wo85a07.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2462100 3'
9726	22377	35578	0.79	1.3E+00	0 AJ223962.1	NT	Lactococcus lactis cremoris NCDO-inv1 chromosomal inversion junction DNA
9726	22377	35579	0.79	1.3E+00		NT	Lactococcus lactis cremoris NCDO-inv1 chromosomal inversion junction DNA
9266	22417	35624	4.53	1.3E+00	1.3E+00 BE963379.2	EST_HUMAN	601657145R1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3866195 3'
9826	22477		0.48	1.3E+00	1.3E+00 AI559944.1	EST_HUMAN	1477412.x1 NCI_CGAP_Uff Homo saplens cDNA clone IMAGE:2214814 3' similar to gb:X14723 CLUSTERIN PRECURSOR (HUMAN);
10050	22698	35913	0.46	1.3E+00	1.3E+00 AF081251.1	LZ.	Eschericia coli serotype 0157:H7 O antigen gene cluster
뎡	22698	35914	0.46	1.3E+00	1.3E+00 AF061251.1	NT	Eschericia coli serotype O157:H7 O antigen gene cluster
10113	22761	35974	1.62	1.3E+00		NT	White chelerae chromosome II, section 49 of 83 of the complete chromosome
g	22778	35991	1.35	1.3E+00		NT	Campylobacter Jejuni kanamycin phosphotransferase (aphA-7) gene, complete cds
10483	23129		0.82	1.3E+00	1.3E+00 AL163302.2	NT	Homo sapiens chromosome 21 segment HS21C102
F	23157	36383	0.45	1.3E+00	6.1	EST_HUMAN	ws32e16x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2498922 3' similar to SW:TRXB_HUMAN Q16881 THIOREDOXIN REDUCTASE ;
10592	23286		4.6			1	DIHYDROPYRIMIDINASE (DHPASE) (HYDANTOINASE) (DHP)

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	1	T	Ţ	T	Ţ	T	T	Т	T	T	T	T	Ţ	T	7	Ţ	T	T	Ţ	T	T	Т	Т	Т	T	Т	Т	Т	T	Т	Т	Т	Т	Т	-
Single Exon Probes Expressed in Brain	Top Hit Descriptor	MRNA 3'-END PROCESSING PROTEIN RNA 45	Mus musculus desmin gene	xp09e03.x1 NCI CGAP HN9 Homo saniens cDN4 clans M44.0F:2730988.9	Human mRNA for KIAA0085 gang, partial cda	Bacillus subtilis genomic DNA 23.9kB fragment	Cavia porcellus inwardiv-rectifung potessium channel Kiro 2 (KCN 112) gene gemeine ede	1802023185F1 NCI CGAP Bm67 Homo sepiens cDNA clone IMAGE-4458457 8	E1 GLYCOPROTEIN PRECURSOR //MATRIX GI YCOPROTEIN / MEMBBANE CI YCOBBATEIN	Sturnira Illium cytochrome b gene, complete ads. milochandrial gene for milochandrial graduat	2/22d08.s1 Soares fetal liver spleen 1NFLS S1 Homo sapiens cDNA clone IMAGE 231838 2	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PEHRP-III)	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PEHRP.III)	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PEHRALIII)	Homo saplens hypothetical protein PRO3077 (PRO3077) mRNA	Elaeis oleifera sesquitarpene synthase mRNA complete oda	pea seed-borne mosaic virus complete genome	pas seed-borne mosaic virus complete genome	Homo sapiens G-protein coupled recentor 14 (GPR14) gene complete and	Homo sablens mRNA for KIAA0874 broken partial cds	Arabidopsis thaliana DNA chromosome 4 contin francent No. 63	Arabidopsis thaliana DNA chromosome 4. contin fragment No. 63	CONJUGAL TRANSFER PROTEIN TRBE PRECURSOR	Mus musculus subtilish-like serine protease LPC (PC7) gene, exons 1 to 9, partial cals	MRo-FT0175-050900-203-g06_1 FT0175 Homo saplens cDNA	Homo sapiens LHX3 gene, Intron 2	Rattus rattus cardlac AE3 gene, excns 1-23	Arabidopsis thaliana DNA chromosome 4, contra fragment No. 21	Homo sapiens post-synaptic density 95 (DLG4) gene. complete cds	T. plnnatum chloropiast rbcL gene, partial	Human extracellular calcium-sensing receptor mRNA, complete cds	MR3-ST0191-140200-013-c05 ST0191 Homo sapiens cDNA	Homo sapiens zinc finger protein ZNF191 (ZNF191) gene. complete cds.	D.hydei ay1 repeat cluster DNA, fragment D	QV4-BN0090-270400-190-a03 BN0090 Homo saplens cDNA
gie Exon Pro	Top Hit Database Source	SWISSPROT	LN	EST HUMAN		Ę	Ę	EST HUMAN		Z.	EST HUMAN	SWISSPROT	SWISSPROT	SWISSPROT	F	ĮN.	LN	LZ	NT.	LN	LN	NT	SWISSPROT	17	EST_HUMAN	TN	F	L	F	N.	LN	EST_HUMAN	Z		EST_HUMAN
uio	Top Hit Acession No.	+00 P25299	+00 Z18892.2	+00 AW274791.1	+00 D42042.1	+00 Z98682.1	+00 AF187873.1		+00 P33464	+00 AF187035.1	H00 AA676246.1		+00 P05228		8924234	+00 AF080245.2		Γ		Γ						.1				1.2E+00 Y09200.1			2.1		
	Most Similar (Top) Hit BLAST E Value	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.3E+00	1.2E+00	1.2€+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00/	1.2E+00/	1.2E+00/	1.2E+00 /	1.2E+00 /	1.2E+00/	1.2E+00	1.2E+00 P54910	1.2E+00 U75902.1	1.2E+00 E	1.2E+00 A	1.2E+00 M87060.1	1.2E+00	1.2E+00	1.2E+00 Y	1.2E+00 U	1.2E+00 A	1.2E+00 A	1.2E+00 X74885.1	1.2E+00 B
	Expression Signal	1.93	2.01	1.8	3.21	3.16	2.64	6.3	2.73	2,15	11.05	0.87	0.87	0.87	1.35	5.64	1.28	1.28	1.22	1.24	96.9	5.98	2.59	69.9	1.78	1.11	1.57	96.0	1.89	6.43	1.1	1.91	0.83	2.51	4.42
	ORF SEQ ID NO:	36744	36772		37433	37544		31035			26050	26239	26240	26241		26554	28600	26601	27463	28512	28573	28574		29091	29354	28766		29850	29892		30836	31158	31504	31801	31865
	SEQ ID NO:	23505	23528	23966	24127		24675	24780	25153	24848	13414	13576	13576	13576	13627	13883	13935	13935	14739	15873	15926	15926	16041	16452	16716	16110	17174	1722	17258	17283	18154	18266	18575	18840	18897
	Probe SEQ ID NO:	10822	10846	11307	11527	11624	12210	12386	12397	12489	932	8	804	804	858	1138	1183	1183	2003	3108	3163	3163	3280	3696	3867	4266	4438	4887	4523	4548	5351	5467	5784	0909	9419

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Single Exon Probes Expressed in Drain	Most Similar (Top Hit Acession Database BLAST E No. Source Value	1 2E+00 X89084 1	-N	1.ZE+00 Appendent	NO INCIDIO	goli/87935jHUMAALU472 Human carcinoma cell-derived Alu RNA transcript, (rRNA); gb:J04970	O RE 1 2F+00 N33285.1 EST HUMAN (CARBOXYPEPTIDASE M PRECURSOR (HUMAN);	1 2E LON P17671 SWISSPROT	4 DE LOOI AWR13276 1 EST HUMAN	1.ZE-500 AVIOLOZI C.:	1.2E+00 AD025010.1	1.2E+00 AJU02141.1	IN TOTAL	S.1 EST HUMAN	1.2E+00 X74207.1 NT	N	1.2E+00 BE787646.1 EST_HUMAN	1.2E+00 AB033030.1 NT		SWISSPROT	1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	NAMULT TO A PASSAGE TO THE MAN	1.2E+00 AWS/12.10.1	1.2E+00 H465995.1	1.2E+00 232850.1	1.ZE+00 D 11.45.	1.2E+00 Aboose 1	1.2E+00 ABOUSSOS.1	1.2E+00 AW 81/81/.1	1.2E+00 BE160761.1	1.2E+00 U50147.1	1.2E+00/AL163203.2	2.8 1.2E+00 AP001515.1 INI	1.53 1.1E+00 D86980.1 INI	1.33 1.1E+00 AW 885385.1 EST III MAN	0.98 1.1E+00 AW5/3889.1 531_10W2/	
-		1 2E+00 X8908	4 7E-00 Year	1.ZE+00 \nood	1.2E+00 AA/5	-	1 2F+00 N332	1 2E 100 P176	4 DE LOO AWR	1.2E+00 APV	1.2E+00 ABV2	1.2E+00 AJ00	1.2E+00 AJ27	1.2E+00 AV73	1.2E+00 X742	1.2E+00 J052	1.2E+00 BE78	1.2E+00 AB03		7 00.70	1 25100	1.25.00	1.2E+00 AW	1.2E+00 F146	1.2E+00 232	1.25-400 011	1.2E+UU A30	1.2E+00 AB0	1.2E+00 AW	1.2E+00 BE1	1.2E+00 U50	1.2E+00 ALT	1.2E+00 APC	1.1E+00 D8c	1.1E+00 AW	1.1E+001AW	
	<u>-</u>	7,87	5 1	7.7	39.54		ם אם	300	0000	2.00	1.17	3.11	0.94	4.86	2.49	0.56	0.56	3.32	-		0.08	10.0	1.87							10.62		=					
	ORF SEQ ID NO:		31951	31952	31989		70700	32701	321/5	32179	32498	32512		33044	L		١			_	34391						35684		37314	2		7 30817		7 25876			
	Exan SEQ ID NO:		18974	18974	19015			19112	19177	19180	19476	19490	19840	1	1	1	١	1				21467			H	22158	22482	22872	3 24009	_	5 23202	9 25227	9 24667	1 13237	7 14499	14629	
	Probe SEQ ID NO:		6198	6198	6241			6342	6408	6412	6815	6829	7153	7282	7550	26037	114	01//	ğ		8561	8775	8923	9138	9298	9906	9831	10224	11318	1135	11435	12179	12199	451	1757	1892	

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Table 4
Single Exon Perhame Expensed

Single Exon Probes Expressed in Brain	Top Hit Descriptor Top Hit Descriptor	NT Homo sanisms chromosome 24 comment 1004.000	- LV	22641 NT	w/64h11.x1 Soares_NFL_T_GBC_S1 Home sapiens GDNA clone IMAGE:2359461 3' similar to	L	Γ	INT						78530 NT	EST HUMAN	EST_HUMAN				HUMAN	N Mus musculus mRNA for ER protein 68 (EP68 gene)			VT Hennes simplex views by to 1 (see III NOS) UL41 gene	NT Arabidoosis Insilana DNA chomosomo 4 actic 10.		H IMAN	Т	NT Acetabularia caliculus mitochandria (COX) Illic and COX (COX)	VH≐anti-cytomegalovirus ghycoprotein B antibody 4D4 heavy chain variable region [human, mRNA Partial, 375] nt]
Single Exon Probes Expre	Top Hit Acession Database Source	Į.	- LV	22641 NT	EST HUMAN	L	N	INT	835331 NT	¥	Į.	LN	6680080 NT Homo seplen	78530 NT	EST HUMAN	EST_HUMAN	11419739 NT Homo senien		Ŋ	EST_HUMAN	N	N N	Į.	LV.	Į.	TN 0967	EST HIMAN	EST HUMAN	TN	Ę
-	<u> </u>	+00 AL 163213.2	+00 AL 163213.2	ş	+00 AI808360.1	+00 AE003886.1	HO0 AE003886.1	+00 XB5374.1	9	30 U3499	+00 U18466.1	00 AJ271740.1	90	о О	1.1E+00 BE980184.1	00 AI138582.1	8		00 AF197861.1	00 KU6037.1	1.1E+00 AJ404004.1	1.1E+00 X55981 1	1.1E+00 Z72338 1	30 Z72338.1	00 AL161588.2		BF6939	1.1E+00 Al478339.1	1.1E+00 AB003088.1	1.1E+00 S80750.1
	Most Simi (Top) Hi BLAST E Value	1.1	1.1E	1.1E	1.1E+	1.1	1.16+	1.16+(1.1E+(1.1E+(1.1E+(1.1E+0	1.15+0	1.11.5	1.1E+0	1.16+0		1.16+0	1.1	4	1.16+0	1.1E+0	1.16+0(1.1E+0(1.1E+00	1.15+00	1.16+00	1.1E+00	1.1E+00
	Expression Signal	6.48	6.48	1.11	1.01	1.05	1,05	1.02	5.69	0.81	3.45	1.05	1.07	1.39	15.75	12	1.		0.62	0.02	0.72	0.72	2.18	2.18	8.84	0.8	3.01	0.64	0.71	0.75
	ORF SEQ ID NO:	28734	28735	28892	28970	29098	29089				30272	30273	30464	30657	31225	31250	31743	00070	31832	32304	1000	32956	33146	33147	33172	33247	33860	33950	34471	34549
	Exen SEQ ID NO:		16084	16236	16322	16460	16460	16550	16831	7369	17662	- 7883	1,847	18031	18324	18342	18782	0000	1000	19379	19842	19882	20068	20068	20088	25115	20727	20814	21328	21406
	Probe SEQ ID NO:	3324	3324	3480	3567	3707	3707	3798	4190	4634	4834	4935	2129	6224	5528	5545	9001	9	8313	8818	7155	7196	7389	7389	7411	7480	8032	8120	8836	8714

Page 25 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	22 ARTE VI SARIAS NHHMPU S1 Homo sapiens cDNA clone IMAGE:1677249 3'	AN 278278F1 NIH MGC 20 Homo saplens dDNA clone IMAGE:3817418 5	Mus musculus mRNA for stretch responsive muscle (X-chromosome) protein (Smix gene)	Arabidopsis thaliana DNA, 24 kb surrounding PFL locus	Yersinia pseudotuberculosis psaE, psaE, adhesin (psaA), chaperone (psab), and using (psab), and using (psab),	complete cds Home sapiens mRNA for KIAA0934 protein, partial cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27	Mus musculus guanine nucleotide binding protein (G protein), gamma 3 subunii (Grigo), mi woo	DNA MISMATCH REPAIR PROTEIN MUTS	151c11.11 Schneider fetal brain 00004 Homo sapiens colors introduced to the colors in the colors of	Human mRNA for 80K-L protein, complete cds. (HUMAN); Homo sepiens KIAA0828 gene product (KIAA0826), mRNA	Klehsormidium fluitans cytochrome c oxidase subunit 2 (cox2) gene, mitochondrial gene encouling	mitochondrial protein, partial cds	domo sapiens potassium inwardly-recurying channel, suggesting of memory of memory of the control	Homo sapiens cytochrome P4502C9 (CYP2C9) gene, 5 flank and exult 1	Homo sapiens hypothetical protein FLJ11280 (FLJ11280), mixiva	Somplete cds	Petroseilnum crispum cytosolic glucose-8-phosphate denydrogenasa T (Coor 2017)	Petroselinum crispum cytosolic glucose & phosphate dehydrogenase 1 (cG6PDH1) mkNA, comprete cos	wf78e11.x1 Soares_NFL_1_GBC_3 I notice September 20.00	Human PBI gene, complete cds	Human PBI gene, complete cas	LOW TEMPERATURE ESSENTIAL PROTEIN	Taenia solium Immunogenic protein 1370 (minora) parasa (Diol) mRNA, complete cds	Dictyostellum discoideum isopenteny pyropriospriate isomecae (Cre), mRNA	Rattus norvegicus C-reactive protein, member of the perfusivity (C.F.)	Xenopus laevis rhodopsin gene, complete cds	Cavia cobaya mRNA for serine/threoline kinase, complete cus	Marchantia polymorpha genes for 26S rRNA, 5S rRNA, 18S rRNA, 3.33 marchantia	Girardia tigrina mRNA for homeodomain transcription factor (so gene)	Homo sapiens chromosome 21 segment HS21 C018	
-	Top Hit Database Source	Т	Т	ES TOWN			-			ISSPROT	Т	T HUMAN		L N				2	본	N	EST HUMAN	LΝ	Į.	SWISSPROT	LN	LN.	LNIS	LN	Į.	NT	Į.	LN L	
	Top Hit Acession No.								1.1E+00/AL161515.2	0704021	13/08	AI87892	11067364 NI	AFORSO42.1	14430508 NT	112909	1.1E+00 L100//.1	887788	+00 AF012862.1	+00 AF012862.1	+00 A1809699.1	+00 D89501.1	4 4E -00 DR9501 1	4 1F+00 P07866	4 1E+00 AF216696.1	4 4E±00 AE234169.1	R393196 NT	1122008 4	1.0E+00 023000.1	1.0E+00 D00423.1	AB02 1004.1	+00 AJZ31000.1	חבומבומב
	B +	Value	1.1E+00 AI079946.1	1.1E+00B	1.1E+00	1.15+00 112227.1	1.1E+00 L76301.1	1.1E+00	1.1E+00/	1.1E+00	1.15+001=73708	1.1E+00	1.1E+00	0	1.15	1.15+00	1.15+00	1.1E+00	1.1E	7	i i	-	1	L								1.0	
	Expression Signal		0.45	0.69	0.53	1.2	1.14	1.37	4.59	18.34	=	0.73	2.25		J.,	1.28	1.58	5.23	3.68			4.30			3.90		2.09	1.44			2.25		4.38
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Table 4

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Most Similar (Top Hit Acession Database BLAST E No. Source Value	1.09 1.0E+00 S52770.1 NT insulin-like growth factor-binding protein 4 (cattle, pulmonary artery endothelial cells, n B-CELL RECEPTOR CD22 PRECURSOR (LEU-14) (B-LYMPHOCYTE CELL ADH	1.0E+00 P20273 SWISSPROT	1.0E+00 AF192631.1 INI	1.0E+00 AA775191.1	1.0E+00 BE808207.1	1.0E+00 BE88820/.1	1.0E+00 D10862.1	4 0E+00 002207 SWISSPROT		1.0E+00 Q02207 SWISSPROT		Т		SWISSPRO!		1.0E+00 Q9Y5T5 SWISSPROT	1.0E+00 BE147331.1 ES TOMEN	Vif protein (vif), Vpr protein (vpr), Tat protein (tat), Rev protein (rev), vpu protein (vpr),	1.0E+00 U42720.2 NT	1.0E+00 M38427.1	1.0E+00 BE907592.1 ESI_HUMAN	1.0E+00 0752429 NT	1.0E+00 6/35428 NI HAAN	1.0E+00 AV68904:1	1.0E+00 044802.1	
		1.0E+00 P20273	1.0E+00 AF1925	1.0E+00 AA7751	1.0E+00 BE8082	1.0E+00 BE8082	1.0E+00 D10852	4 0F+00[00220]	20. 10.	1.05+00 00220	31	1.0E+00 P5178		1.0E+00 Q9Y5T		1.0E+00 Q9Y51	1.0E+00 BE147	<u> </u>	1.0E+00 U4272	1.0E+00 M384	1.0E+00 BE90	1.0E+00	1.0E+00	1.0E+00 AV68	1.0E+00104+30	1,00,000
Expression (Signal E	1.09	9.29	1.56	5.26	1.36	1.36	1.19					0.85												ļ		
ORF SEQ ID NO:	32788		33385						١								Ш									39 35352
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	Exan ORF SEQ Expression (Top) Hit Top Hit Acession Database SEQ ID NO: Signal Value No.	Exon ORF SEQ Expression (Top) Hit Sequences (Top) Hit No. Top Hit Acession (Top) Hit Top Hit Acession (Top) Hit No. Top Hit Acession (Top) Hit Top Hit Acession (Top) Hit No. Top Hit Acession (Top) Hit Top Hit Top Hit Acession (Top) Hit Top Hit	Exon SEQ ID NO: ORF SEQ Signal NO: Expression Signal Value Most Similar (Top) Hit No. Top Hit No. Top Hit Database Source Value 19730 327788 1.09 1.0E+00 S52770.1 NT	Exon SEQ ID NO: ORF SEQ Signal ID NO: Expression Signal Signal Signal NO: Most Similar Top Hit No. Top Hit No. Top Hit Source Source Value 19730 32788 1.09 1.0E+00 S52770.1 NT 20058 9.29 1.0E+00 P20273 SWISSPROT SWISSPROT FEST HIMAN	Exon No: ORF SEQ ID ID NO: Expression Signal No: Top Hit Acession (Top) Hit Top Hit Acession Source No: Top Hit Acession Source No: Top Hit Acession Source Naive Top Hit Acession Source Naive Top Hit Acession Source NO: 32788 1.09 1.0E+00 S52770.1 NT 20058 9.29 1.0E+00 P20273 SWISSPROT 20277 33386 1.66 1.0E+00 AF192831.1 NT 20277 5.26 1.0E+00 AF715191.1 EST HUMAN 20287 33401 5.26 1.0E+00 AF775191.1 EST HUMAN	Exon ORF SEQ Expression Signal Top Hit Acession Top Hit Top Hit Acession Source Top Hit Top Hit Acession Source NO: Signal Signal 1.09	Exon ORF SEQ Expression Signal (Top) Hit Acession Value Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Source NO: Signal 1.09 1.0E+00 S52770.1 NT I 20058 9.29 1.0E+00 P20273 SWISSPROT 20277 33385 1.66 1.0E+00 AF17519.1 EST_HUMAN 20556 33401 5.26 1.0E+00 BE868267.1 EST_HUMAN 20556 33681 1.36 1.0E+00 BE868267.1 EST_HUMAN 20556 33682 1.36 1.0E+00 BE868267.1 EST_HUMAN 20556 33682 1.36 1.0E+00 BE868267.1 EST_HUMAN 20556 33682 1.36 1.0E+00 BE868267.1 EST_HUMAN	Exam ORF SEQ Expression Signal (Top) Hit Accession No. Top Hit Accession Source Top Hit Accession Source NO: Signal Signal Signal No. 1.0E+00 S52770.1 NT II 19730 32788 1.09 1.0E+00 S52770.1 NT II 20058 9.29 1.0E+00 P20273 SWISSPROT SWI	Exam ORF SEQ Expression Signal (Top) Hit Acession No. Top Hit Acession Database Source Source Source Top Hit Acession <	Exam ORF SEC Signal Expression Top) Hit Value Top Hit Acession No. Top Hit Acession Source To	Exon No: ORF SEQ Signal Signal Most Similar Top Hit Acession Source Value Top Hit Acession Database Source Source Value Top Hit Acession Database Source Source Value NO: 32788 1.09 1.0E+00 S52770.1 NT ii 20058 9.29 1.0E+00 P20273 SWISSPROT (PEST HUMAN FIGURES).1 NT H 20277 33386 1.66 1.0E+00 A775191.1 EST HUMAN FIGURES EST HUMAN FIGURES 20566 33682 1.36 1.0E+00 BE868267.1 EST HUMAN FIGURES 17680 1.0E+00 BE868267.1 EST HUMAN FIGURES SWISSPROT FIGURES 17680 2.05 1.0E+00 BE868267.1 EST HUMAN FIGURES 17680 2.02 1.0E+00 BE868267.1 EST HUMAN FIGURES 17680 2.02 1.0E+00 BE868267.1 EST HUMAN FIGURES 17690 2.02 1.0E+00 BE868267.1 EST HUMAN FIGURES 17690 2.02 1.0E+00 BE868267.1 SWISSPROT FIGURES	Exon No: ORF SEQ Signal Signal Most Similar Top Hit Acession 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Exon ORF SEQ Expression Top Hit Top Hit Acession Political Polit	bes Expressed in Brain	Top Hit Descriptor		Human Coronavirus gene for membrane protein	Human Coronavirus gene for membrane protein	Homo sapiens MHC binding factor, beta (MHCBFB) mRNA	Homo sapiens MHC binding factor, beta (MHCBFB) mRNA	oy15407.s1 Soares, senescent fibroblasts_NbHSF Homo saplens cDNA clone MAAGE-4885004 pi	AV768825 BM Homo septens cDNA clone BMFAWC04 5	Zn94a02.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE 42860a e/	ZIB4802.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGF-4280ng F:	Infilman retnoblastoma susceptibility gene exons 1-27, complete cds	Por I-Digine-field protein (Infron 3) [human, Genomic, 898 nt] 263b11.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo saniens cDNs_chan_1140CF_13555555	contains Alu repetitive element; contains element MER38 repetitive element:	Picea glauca EMB13 mRNA	Hordeum vulgare gene encoding cysteine proteinase	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 8 (NADH DEHYDROGENASE 1, CHAIN 8) ANDU 1	(Internal of the control of the cont	Human adenovirus type 5, complete genome	I HROMBOMODULIN PRECURSOR (FETOMODULIN) (TM)	EST 388293 MAGE resequences, MAGN Hamo sapiens cDNA	rigino saplens chromosome 21 segment HS21C102	Apple mosalc virus RNA 2 putative polymerase gene, complete cds	SENING! HKEUNINE PROTEIN KINASE MINIBRAIN	NOBABLE OXIDOREDUCTASE ZK1280.5 IN CHROMOSOME II	-ycopet sicon esculentum putative Mi1 copy 1 nematode-resistance gene	SE ENAUTININ RECEPTOR (BK-2 RECEPTOR)	Janio rerio mKNA for Eph-like receptor tyrosine kinase rtk8	Saprificola 16S rDNA (host T.suberl)	.aphidicala 16S rDNA (host T.suberi)	IMINO-ACID ACETYLTRANSFERASE (N-ACETYLGLUTAMATE SYNTHASE) (AGS) (NAGS)	Allithrix Jacchus UBE1 gene derived retroposon on the Y chromosome	enopus laevis rac GTPase mRNA, complete cds	incorbere ENDONUCLEASE IV (ENDODEOXYRIBONUCLEASE IV) interobacteriaceae sp. JM983 partial groES gene for GroES-like protein and neutral groES gene for GroES-like protein and neutral groES.	Ce protein, isolate JM983
E EXON ORF SEQ Expression ID NO: Most Similar Similar Signal Most Similar No. No. 32 22404 35609 0.49 1.0E+00 X15498.1 No. 32 22404 35609 0.49 1.0E+00 X15498.1 2 2260 36875 0.71 1.0E+00 X15498.1 2 2260 36876 0.71 1.0E+00 X15498.1 4 22762 36985 0.71 1.0E+00 X15498.1 5 23021 36237 1.0E+00 AND40482.1 6174 5 23021 36237 1.1 1.0E+00 AND40482.1 5 23021 36237 1.1 1.0E+00 AND40482.1 5 23021 36237 1.1 1.0E+00 AND40482.1 5 23021 36239 1.1 1.0E+00 AND40482.1 5 23021 37623 1.1 1.0E+00 AND40482.1 1 8008 1.59 1.0E+00 AND40484.1 2 24122 37794 1.38 1.0E+00 AND40484.1	ngle Exon Pr			Z	Z	N	- In	EST HUMAN	EST HUMAN	EST HUMAN	ES L HUMAN	L L		EST HUMAN	Z	2	TOGGGG	TO LOCAL DE LA COLONIA DE LA C	1000000	SWISSPRO	TOWN TO	1	Τ.	Т	Т	┰				100000	ISSPEC		TOBOS!		
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Exon ORF SEQ Express		Most Similar (Top) Hit BLAST E Value	1 OF ±00	105+50	1 OF +00	10.1	100-100	10F+00	105+00	105+00	1.0F±00	1.0E+00	1 08100	1 05-400	1 OF +OO		1.0E+0010	1 05+00	1 OF +OO	1.0E+00	9.9E-01 A	9 9F-01 A	9.9E-01 P	9.9E-01 Q	9.9E-01 U	9.9E-01 Q	9.9E-01 A.	9.9E-01 Y	9.9E-01 V	9.8F-01 P	985-014	9.8F-01 A	9.8E-01 00	9.8F-01 A.1	
SEQ 1D OR		Expression Signal	0.49	0.49	0.71	077	0.81	4.36	16.18	16.18	-	4.57	1 48	1.59	1.55		12.29	1.38	3.04	2.32	1.19	0.97	10.09	0.93	1.37	2.18	2.37	2.3	2.3	1.14	121	1.01	0.92	4.67	
0 888888888888888888888888888888888888			L				L				36270	36823	38960		30629		37763	37794	-		28097		31253	31498			36547	37508	37509	25926		-	29171	32852	
Probe SEQ ID NO: NO: 10012 10012 10012 10012 10012 10012 11622 11622 116614 116			L		LI		L						23697	24122	18008		24422	24452	24588	24772	15353	18345	18344	18570	21830	22005	23308	24191	24191	13294	15020	15509	16533	19788	
		Probe SEQ ID NO:	9763	9753	10012	10012	10104	10225	10375	10375	10407	10893	11025	11522	11744		11838	11868	12049	12370	2643	3591	5547	5779	9160	9455	10614	11592	11592	510	2295	, 2804	3781	7099	

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	Top Hit Descriptor Top Hit Descriptor	Enterobacteriaceae sp. JM983 partal grocs gene for order processing processin	0014503011 1711 BHOMO sepiens cDNA clone IMAGE:3830049 5	PHOSPHOGLUCOMUTASE (GLUCOSE PHOSPHOMUTASE) (PGM)		601110258F1 NIH MGC 16 Homo saplens cDNA clone IMAGE:3350750 5	601110258F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3330730 3	042c10.x1 NOI_CGAP_Lu24 Homo saplens cDNA done IIV/AGE_2212222	Homo sapiens X28 region near ALD locus containing dual specificity prospiratives of U.O.O. 7, 11200000000000000000000000000000000000	protein L18a (RPL18a), Ce2+/Calmodulin-dependent protein furescriptory. CDM protein (CDM), adrenoleukodystrophy protein CDM protein (CDM), adrenoleukodystrophy protein (CDM), adrenoleukodystrophy protein (CDM).	Drosophila meanogaster boulan organic promiter of a partial cds	Triticum aestivum stripe rust resistance protein Y110 (1710) gane, without ond principassa (res)	Salmonella typhimurium adenine-methyltransferase (mod) and resurction and unward (100)	UI-H-BI4-aci-e-07-0-UI.s1 NCI_CGAP_Subs Homo saplens cDNA clone IMACE. 3050-1-3	PM2-UM0053-240300-005-f12 UM0053 Homo saplens cUNA	Parvovirus B19 DNA, patient C, genome position 2446-2994	Parvovirus B19 DNA, patient C, genome position 2448-2994	Arabidopsis thaliana DNA chromosome 4, ESCA I TON COUNTY inspirior	P falciparum complete gene map of plastid-tike DNA (IN-A)	Rattus norvegicus (strain R21) Rpszt gene, comprese cus	AV752805 NPD Homo sapiens cUNA clone NPDBAG06 5	AV752805 NPD Hamo suplens control material of (CEP2) mRNA	Homo sapiens centroscaria process are processed into the processed int	Sphyma doulo NADh deugal againeachail againeacha protein, partal deugan an ann	Homo saptens CGI-129 protein (COC) (ENDOGLUCANASE I PRECURSOR (EGI) (ENDOT), TOTAL MAGE: 3958473 5	601675639F1 NIH_MGC_21 Homo saplens cDNA clone invo E-secard R	601675639F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE: 3304173581 3/	qd57d07.x1 Soares_testis_NHT Home sapiens cultar cruite ilwaycer	RC1-CT0295-241199-011-b02 CT0295 Homo sapiens culvin	
Single Exoli Flores	Top Hit Detabase Source	\Box	Т	EST HOMAIN	\neg	EST HIMAN 6	Т	Т	Г	ž.	<u> </u>			HIMAN	HUMAN	Т					THUMAN	HUMAN	FN	NT	LN	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	
Buio	Top Hit Acession No.			9.1	١	١			UI Alconovic.	U52111.2		,		1	T	9.6E-01 AW 799074.1		9.0E-01 £703341 2	9.0E-01 28/341.2	04 84438 4	9.9E-01 AV752605.1	AV7526	11421722 NT	U91423.1	7705591 NT	002834	8.05-01 Q02001	9.5E-01 BE902340.1	9.3E-01 DE90203	9.5E-01 A1180 102.1	2000
	Most Similar (Top) Hit BLAST E Value	9.8E-01 A	9.8E-01 B	9.8E-01 B	9.8E-01 P	9.8E-01 A	9.8E-01 BE258/05.1	9.8E-U1	9.85-01	9.8E-01	Z L	9.7E-01 0207 10.1	9.7E-01	9.7E-01 M90544.1	9.7E-01	9.65-01	9.6E-01 Z/0556.1	8.00	9.00	8.00-0	9.0F.0	9.6E-01	9.6E-01	9.6E-01	0 5E-01	0.00 1	8.05.0	9.55-01			
	Expression (4.67	1.15	1.15	0.91	1.13	2.29	2.29	1.57	1.56		2,28	1.68	1.3	3.64	1.5	3.77	3.77	0.61	2.3	0.08	3.47	1.92	21.0	2 7	1.03					1.05
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Table 4
Single Exon Probas Everand

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		601885163F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGF-4103830 F	UI-H-BI2-ahp-f-03-0-UI.s1 NCI_CGAP_Sub4 Homo sapiens cDNA close IMADE:0727222 2.	ya53d04.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMA GE-8834 of	Bartonella claritigeiae RNA polymerase beta subunit (rooB) dene partial car	Pimpinella brachycarpa zinc finger protein (ZFP1) mRNA complete ede	Human Fc-gamma-receptorIIA (FCGR2A) gene, exon 4	601466703F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3869929 67 Homo saplens epidermal growth factor recentive (when parties is a second parties of the part	homolog) (EGFR), mRNA	Homo saplens phytanoyi-CoA hydroxylase (PHYHI) gene exp. 5	RC5-BT0503-271199-011-B01 BT0503 Home seniers CDMA	Bovine papillomavirus type 2, complete genome	Bovine papillomavirus type 2. complete genome	Homo saplens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NEKB1)	cas	Spodobera frugiperda methylenetatrahydrofolate dehydrogenase mRNA commissioned	0903903.s1 NC_CGAP_Ovz Homo sepiens cDNA clone IMAGE:1385357	Xenopus laevis CCCH zinc finger protein C3H-2 (C3H-2) mRNA, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 34	Aedes triseriatus putative large subunit ribosomal protein rpl.34 mRNA, comulate ode	Homo sapiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A, melanoma antigen family A12 (MAGEA12), melanoma antigen family A2b (MAGEA2B), melanoma antigen family A2b (MAGEAB), mel	(CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and Lis and Lis	501441338T1 NIH_MGC_72 Homo saplens cDNA clone IMAGE 3016184 3	601817814F1 NIH_MGC_58 Homo saplens cDNA clone IMAGE:4041363 F/	Mus musculus solute carrier family 30 (zinc transporter), member 4 (Sic30e4), menta	601461153F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE 3844641 E	N.crassa valyi-tRNA synthetase (cyt-20/un-3) gene	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 65	Mus musculus carbonic anhydrase 4 (Car4) mRNA	Homo saplens lysosomal apyrase-like protein 1 (LALP1), mRNA	Podeso Na NCI CGAP XId11 Home sapiens cDNA clone IMAGE:3578219 3' similar to SW NUSM TRYRR	801334943E1 NIH MGC 30 HCC 20	Committee of the commit
ngle Exon Pr	Top Hit Database Source		EST HUMAN	EST HUMAN	ESI HUMAN	Z	L _N	LZ.	EST_HUMAN	NT	LN	EST_HUMAN	Į,	N	Ļ	LIN LIN		ES - HUMAN		FIN			7	7	INMAN		HOMAN					EST HUMAN	7	7
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	Most Similar (Top) Hit BLAST E Value	0 AE 04	9 GE 04	0.5E.04	9.4F-04	0.45.04	9.45	9.45-01	10 1	8.4E-01	9.35-01	9.35-01	8.35-01	9.3E-01	9.3E-01	9.3E-01	935-010	9.3E-01	9.3E-01 A	9.3E-01 A		9.3F_04	9 2E 01 B	0.2E.04 BE420072 4	9 27.01	9.2F_01 RE037598 4	0 2F-01 MARAZO 4	0.0E.04 Al 40450F.0	0 2E 04	100	9.45-01	9.2E-01 BF		
	Expression Signal	1,68	2.42	1.55	3.33	2.08	0 87	1 92	,	100	1.38	8 0	3 6	0.08	1.56	3.89	1.85	1.04	0.95	3.12		1.48	2 93	26.0	1.15	7.36	0.61	0 92	107	2 18	2 5	1.9	1.63	
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	Probe SEQ ID NO:	11209	11429	11795	3196	3212	8764	12202	12557	1726	2640	4015	4015		5505	2293	7966	8713	8835	12881		12802	3233	4822	5631	2888	6537	9560	9648	10165	i	10315		
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| Top Hit Descriptor | nn05f11.s1 NCI CGAP Pr4.1 Home septems CDNA clone IMAGE 40768277 | Pseudomonas aeruginosa topoisomerase (top), putative transcriptional regulatory protein OhbR (ohbR), ortho-
halobenzoate 1,2-dioxygenase beta-ISP protein OhbA (ohbA), OhbC (ohbC), ortho-halobenzoate 1,2- | Homo septents partial 1 GAT So and Combb, and pure | RCANNOST 170500 043 -073 MILEATT | ob38e08 x1 Source, NEI T ODC Sx U | divided by Source 1 CBC 51 Home septens CDNA clone IMAGE:1846786 3 | Process of States and States and States of States
of States of Sta | 60218544174 NIH MGC 45 L | 80218554111 NIH MGC 48 Line 12 Trough Septemble Colore IMAGE:4309906 3 | OVO-NN1021 100500 227 - 22 Million Baberls CLINA clone IMAGE:4309908 31 | R0182388481 NIU MCC 36 L | GOTA STABLES NIH MICE TO LINE SEPTEMBER CLINA CIONE IMAGE: 4043564 3' | Bet (DEI) 2000 for including a north a spiens a UNA cione IMA(GE:4043564 3)
 | The state of the s | Homo saplens cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebrotendinous

 | xanthomatosis), polypeptide 1 (CYP27A1b) mRNA | Arabidopsis thallana DNA chromosome 4, contig fragment No. 65 | Urosophila melanogaster merlin (Dmerlin) mRNA, complete cds | Chloken lipase gene | Gris renedensis | Grus canadensis recombination political activation of the control of
the control of the control | Helicobacter pylori 28805 seekin 80 of 424 - 64. | Berillis Paleduren consoli DNA | Drosobila melanoraster collensin research | Archaeoglobus fulgidus saction 128 of 172 of the complete cds | Sotrytis cinerea strain T4 cDNA (thrent index conditions)
 | Sacteriophage D3. complete genome
 | 301067107F1 NIH MGC 10 Homo sapiens cDNA close MAGE 2482565 51 | SEGMENTATION PROTEIN PAIRED | SEGMENTATION PROTEIN PAIRED | Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5 |
| Top Hit
Database
Source | EST_HUMAN | F | L. | EST HIMAN | EST HUMAN | EST HIMAN | LN
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| | AA595863.1 | AF121970.1 | AJ288085.1 | AW897335.1 | AI239456.1 | A1239468.1 | AE004963.1
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(Top) Hit
BLAST E
Value | 8.7E-01 | 8.7E-01 | 8.75-01 | 8.7E-01 | 8.7E-01 | 8.7E-01 | 8.7E-01
 | 8.7E-01 | 8.7E-01 | 8.7E-01 | 8.7E-01 | 8.7E-01 | 8.6E-01
 | 8.6F-01 | 19 0

 | 0.05-01 | 8.85-01 | 9.0E-01 | 8.6E-01 | 8.6E-01 / | 8.6E-01
 | 8.6E-01 | 8.6E-01 | 8.6E-01 A | 8.6E-01 | 8.6E-01
 | 8.5E-01 A
 | 8.5E-01 B | 8.5E-01 P | 8.5E-01 P | 8.5E-01 A |
| Expression
Signal | 6.05 | 3.17 | 76.0 | 0.62 | 0.69 | 0.69 | 1.57
 | 0.61 | 0.61 | 5.25 | 5.47 | 5.47 | 1.75
 | 3.45 | 80.0

 | 00.0 | 1 85 | 40 01 | 10.86 | 2.06 | 2.08
 | 0.78 | 1.12 | 0.55 | 0.46 | 1.35
 | 0.95
 | 2.51 | 0.78 | 0.78 | 0.67 |
| ORF SEQ
ID NO: | 28286 | - | | 33761 | 34665 | 34666 | 35483
 | 38065 | 38066 | 36665 | 37657 | 37658 |
 | 26279 | 27733

 | 20002 | 20172 | 31524 | 31526 | 32385 | 32386
 | | | 33763 | |
 | 32401
 | 33189 | 34147 | 34148 | 34231 |
| | Ш | 17673 | | | | . 1 |
 | - 1 | - 1 | 1 | | 24332 | 13247
 | 13608 | 14994

 | 18381 | 16534 | 18597 | 18597 | 19372 | 19372
 | 20104 | 20523 | 20836 | 8223
7223 | 25144
 | 9388
 | 20102 | 21010 | 21010 | CROLZ |
| Probe
SEQ ID
NO: | 2877 | 4946 | 5102 | 7939 | 8828 | 8828 | 9638
 | 10202 | 10202 | 10735 | 11739 | 11739 | 462
 | 838 | 2288

 | 3608 | 3782 | 5808 | 5808 | 6099 | 6099
 | 7427 | 7828 | 7941 | 9586 | 12618
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| | Exon ORF SEQ Expression (Top) Hit Top Hit Acession No. Signal BLASTE No. Source | Exon ORF SEQ ID NO: Expression Signal (Top) Hit Accession Value Top Hit Accession No: Top Hit Accession N | Exon No: Signal Most Similar (Top) Hit Acession NO: Signal Top Hit Acession NO: Signal Top Hit Acession NO: Signal Top Hit Acession NO: Signal Top Hit Acession NO: Signal Top Hit Acession NO: Source No No: Source No NO: Source | Exon ORF SEQ ID Expression Signal (Top) Hit Acession Value Top Hit Acession Signal Top Hit Ac | Exon ORF SEQ ID Expression Signal Most Similar (Top) Hit Accestion Signal Top Hit Accestion No. Top Hit Accession Signal Top Hit Accession Signal Top Hit Accession No. Top Hit Accession Source | Exon ORF SEQ ID Expression Signal Most Similar Top Hit Accession Signal Top Hit Accession No. Top Hit Accession Signal Top Hit Accession Signal | Exon ORF SEQ ID Expression Signal Most Similar Top Hit Accession Signal Top Hit Accession No. Top Hit Accession Signal Top Hit Accession Signal Top Hit Accession Source | Exon NO: ORF SEQ ID NO: Expression Signal Signal Most Similar Top Hit Accession No: Top Hit Accession Signal Top Hit Accession No: Top Hit Acces | Exon ORF SEQ ID NO: Expression Signal Most Similar Top Hit Acession Signal Top Hit Acession No: Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession No: Top Hit Acession Signal Top Hit Acession Signal | Exon NO: ORF SEQ Expression NO: Expression Signal Signal Signal Signal Signal Signal Signal Signal NO: Most Similar No: Top Hit Acession Value No: Top Hit Acession Signal No: Top Hit Acession No: Top Hit Acession Source Source Source Source Source Source Source Source Source Source Source Source Source No: Ace Segon Source
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Most Similar Top Hit Acession Database Top Hit Source Source	DA B006799.1	O CE DA ABORAZOS A INT	8.0E-01 ABOUT 99.1	TIM C STOCOGO A LO TL C	AF083975.2 IN 1	8.4E-01 L/8/20:1	8.4E-01 L/8/20.1	8.4E-01 AF-031142.1 NT	8.4E-01 AJZ46287.	8.3E-01 M93437.1	8.3E-01 AL101500.2	8.3E-01 AB0108/9.1	8.3E-01 Y19177.1	N	T_HUMAN	8.3E-01 AF098070.1 NT	N.	3.35 8.3E-01 AE000903.1 NT complete genome		8.3E-01 AF020503.1 NT	8.2E-01 AB000489.1 NT	8.2E-01 AF145589.1	8.2E-01 AW376990.1 EST_HUMAN	8.2E-01 AJ010142.1	01 AW379433.1 EST HUMAN	01 Z12126.1 NT	8.2E-01 BE263145.1 EST_HUMAN	8.2E-01 AB014530.1 NT	8.2E-01 AF052659.1 NT	0.59 8.2E-01/AF223889.1 NT Olifoting interpretation of the company
	ξ	-		0.35-01	8.4E-01	8.4E-0	8.4E-01	8.4 1.0	1		١					L		l		8.3E.4	8.2E-	8.2E-	8.2E-	8.2E-	8.2E-	8.2E	8.2	8.2E	8.2E	8.2E
Expression Signal	1,17	-	1.17	7.24	0.73	2.28	2.28	0.63	2.68	2.8	2.99	0.79	3.35	2.41	4.51						2		1.0							
ORF SEQ ID NO:	30,00	30100	36108	1	30067	30910	30911	33485		26151		77162		30511		35872				72077				32631			L			l
Exon SEQ ID NO:		22896	22896	25296	17436	25068	25088	20372	22508	13498		١	1	Ĺ	<u> </u>	1	1		78020		L		1	1	ı	i	1	1	1	ı
Probe SEQ ID		10248	10248	12278	4702	5408	5406	7708	8888	724	3091	3790	3993	5187	61.0	8300	10118		7/001	OS S	112/4	300	2686	8878	6797	71.80		7000	a g	

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Table 4
Single Exon Prohas Expensed

Public Exam Cope SEC Cope			_	_	_	_																		٠									
Colored Colo	bes Expressed in Brain			MCKUSICK-KAUFMAN/BARDET-BIEDI SYNDROMES BITATII'IT ALLI	MCKUSICK-KAUFMAN/BARDET-BIEDL SYNDROMES PUTATIVE CHAPERONIN	Molluscum contaglosum virus type 1 ORF1 and ORF2 DNA	OVARIAN TUMOR LOCUS PROTEIN W14402 H Sparse Blooms Profession	similar to gb:M36072 60S RIBOSONAL DECEMBER 11 11 11 11 11 11 11 11 11 11 11 11 11	Mus musculus mRNA for NIPSNAPD mater	Mus musculus TANK binding kinase TBK1 (Tbk1) mBNA	Homo saplens MHC class 1 region	Homo sapiens MHC class 1 region	Drosophila melanogaster Na/K-ATPase beta subunit ischer 47 No. 1-21	Mus musculus putative collegen alpha-2 (XI) chain (COI 1142)	VEURONAL MEMBRANE GLYCOPROTEIN MA. B	VEURONAL MEMBRANE GLYCOPROTEIN MB.B	Jomo saplens KÍAA0417 mRNA, complete cds	SYTOCHROME B	Prosophila melanogaster putative inorganic phosphate cotransporter (Picot) gene, partial cds; putative sodium hannel (Nach) and putative amylase-related protein (Amaria)	nriched protein (gprs) gene, partial cd>	rosophila melanogaster putative inorganic phosphate odransporter (Ploot) gene, partial cds; putative sodium namel (Nach) and putative amylase-related profesh (American)	iniched protein (gprs) gene, partial cd>	acilius halodurans genomic DNA, section 11/14	adillus halodurans genomic DNA, section 11/14	01h03.x1 NOI_CGAP_Kid11 Homo saplens cDNA clone INAGE:2892469 3' similar to SW:LYAR_MOUSE 08288 CELL GROW TH REGULATING NI ICLED AD BEACHER.	ament;	KOBABLE E4 PROTEIN	9872F Human fetal heart, Lambda ZAP Express Homo sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara KKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens CDNA clara kKoozz er aller and a state of the sapiens contains and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er aller and a state of the sapiens clara kKoozz er	(CLONE CAPE11)	20-1 N0080-220800-025-d10 TN0080 Homo sapiens cDNA	July N0080-220800-025-d10 TN0080 Homo sapiens cDNA	amotoga maritima section 23 of 136 of the complete genome	phylococcus aureus partial pta gene for phosphate actyltransferase allele 15
be Exon ORF SEQ ID SEQ ID NO: Expression Signal (Top) Hit Top Hit Acess No. Most Similar No. 283 22931 36145 3.66 8.2E-01 QSJI70 283 22931 36146 3.66 8.2E-01 QSJI70 284 22931 36146 3.66 8.2E-01 QSJI70 284 22933 37659 6.02 8.2E-01 P10383 40 24328 37652 6.38 8.2E-01 P10383 31 24328 37659 6.02 8.2E-01 P10383 40 24333 37659 6.02 8.2E-01 P10383 40 24328 37659 6.02 8.2E-01 P10383 51 16207 28857 3.08 8.1E-01 AF065066.1 51 16207 28867 3.08 8.1E-01 AF065066.1 51 16207 28868 0.74 8.1E-01 AF065066.1 51 16207 28868 0.76 8.1E-01 AF06777.1 52069 33628 0.75 8.1E-01 AF06777.1	gle Exon Pro	Top Hit Database Source		SWISSPROT	SWISSPROT	NT	SWISSPROT	T_HUMAN							٦	ISSPROT	٦	П									Т		T	Т	丁		30
D SEQ ID ORF SEQ Expression ORF SEQ	Sin	Top Hit Acess	1								T											T	I	T		T			-		T	T	7
be Exon ORF SEQ Express 22931 36146		Most Simila (Top) Hit BLAST E Value	ł	ı		2 3 A	0.32.0	8.2E-01	8.25-01	8.1F-01	8 4E 04	8 11 5	8 1E 01	8 1 1 0 1	8 45 04	8 1110	8 1E.04		α T	100	8 1E-01	8.1E-01/	8.1E-01		7	8 15 01 0		8.1E-01	8.1E-01B	8.1E-01 BI			
be Exon NO: 70 N		Expression Signal	3 85	3.65	4.65	6.38		20.02	4.07	3.08	3.08	0.74	0.64	2.66	2.66	0.78	0.65		0.75		0.75	0.93	0.93		108	0.7	-	0.5	2.63	2.63	1.57	3.49	
8 □ 8 8 1 9 9 8 8 1 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		ORF SEQ ID NO:	36145	36146	37562	37652	37650	31054		28857	28858		31973	32295	32296	32987	33173	-	33628	-	33629	34344	34345	_	34507	35888	-	36174	37374	37375	31110		
8 2 2 8 8 8 7 9 9 8 8 8 8 8 8 8 8 8 8 8 8 8 8			Ш		1	li	L	Ĺ		16207	16207	17592	18997	19292	19292	19914	20089	-	20506	_	20506	21199	21199		21360	22673	-	22958	24067	24067	24550	12985	
		Probe SEQ ID NO:	10283	10283	11641	11735	11740	12298	2762	3451	3451	4863	6223	9259	8258	7229	7412		7811		7811	8507	8607					_	┙			_	

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		INTERLEUKIN-6 PRECURSOR (II_A)/P OF I OF II	BETA-2) (HYBRIDOMA GROWTH FACTOR SHIMULATORY FACTOR 2) (BSF-2) (INTERFERENCE	Themoplasma acidonali importation (2007)	164005 xt Screen Not From the Benome; segment 4/5	Didiscolder in record by OT PA P S1 Homo saplens cDNA clone IMAGE 3656472 21	Omo sanlare musicant gene	MUSCIE CALCITIM CHANKING (AIN) (NUP214), mRNA	Arabidopsis thaliana 1-amino-1	Special Complete Comp	Copersicon hirsutum ADP-clucose manhan-1	Mus musculus major histocompatibility locus class II region	chain (Kalpha) and major histocompatibility protein class II had a feet of the class II alpha	typophilin-like (NG9), butyrophilin-li>	CITRATE SYNTHASE	Homo saplens UDP-N-acetyl-alpha-D-galactosamina-nakanaka	ainAc-T7) (GALNAC-T7), mRNA	Homo sapiens PRO1975 mRNA cremulate ada	Columb columb Japonica sub-species language	urnix cofurnix laborice sub-species personal personal managements and services sub-species	RAFFINOSE INVERTASE / INVERTAS	RAFFINOSE INVERTASE (INVERTASE)	1002,s1 Soares fetal liver spleen 1/15 CL	Daphnia magna hemoglobin gene cluster / Ahba Abba	Archaeoglobus fulgidus, complete genome	Arabidopsis thaliana 3-methylcrotonyl-CoA carboxylase non-binthylased anti-mit discourse	Idopsis thelians 2 met. 3	cds	MATING-TYPE PROTEIN 4	51 NOTE IN A-ALPHA 24	and the 2 stell of the septent of th	Rattus noneclarus 11:	Section in independent alpha-latrotoxin receptor mRNA, complete cde	
ingle Exon Pro	Top Hit Database Source			ISSPROT		T_HUMAN			ISSPROT	TN		LN L	2	5 J	7	SWISSPROT								HUMAN						_	EST HUMAN 8014		_		
S	Top Hit Acess No.		8E-01 P05234	1,0000	0E-01 AL 445066.1	oc-01 BF-108927.1	710159.1	4826873 NT	E-01 Q25452	E-01 L29260.1	A 7. 6. 6. 6.	L-01 AF 184345.1		57.1			839340RINT	-01 AF118085 4		T				1	-01 1407824 NT	1170/6111	01 AF059510.1 NT								
	Most Similar (Top) Hit BLAST E Value		7.8E-0	7 85 0	7.05-0	0-30.7	7.85-0	7.8E-01	7.8E-01	/.8E-01	7 75.04			7.7E-01	7.7E-01 033915		7.7E-01	7.7E-01/	7.7E-01.	775.01	20 34.	7 7E-04 D4-015	7 7E 04 0	775.01	7.7E-01	-	7.6E-01 A	100	7 8E 04 PG 250	7.05-01 1-37938	7.8E-01 AI	7.6E-01 AI253399.1	7.8E-01 U72487.1		
	Expression Signal		0.88	0.63	133	100	5 6	10.0	/ACO		7.61			2.26	2.21	-	0.84	3.98	3.38	3.38	145	1.45	0.85	0.51	4.55		4.88	4 88		1	1000	48.0	0.88		
	ORF SEQ ID NO:		31871	32136	34216	34987	35083		1		25596			20,00	28183	·		28985	29738	29739	31165	31166	31594	35600	H	70	10/10	31752	32189	30555	30585	32440	2		
	SEQ ID NO:		4		_	21821		L	L		12954		13/00	15/2/	1750	-	10111	2	7.183	17103	18272	18272	18653	22395	24844	18780	3 2	18789	19193	17920		L			
	Probe SEQ ID NO:	-	47LD	58	8380	9133	9231	10024	12271		139		200	27.17		3254	3588	3 25	202	3	5473	5473	998	9744	12161	8008	+			6751	6751	L	1		

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Single Exon Probes Expressed in Brain		Top Hit Descriptor		Orzies latipes gene for membrane quantify cyclasa OIGC1 complete add	Oryzlas latipes gene for membrane guanyly cyclase OIGC1 complete cds			HYPOTHETICAL PROTEIN HKLF1 (IRL1) (TRL1)	Borrelia burgdorfari (section 52 of 70) of the complete genome	Homo sepiens HT017 mRNA, complete ods	FERRICHROME SIDEROPHORE PEPTIDE SYNTHETASE	Mus musculus antgen (CD72) gene	Mus musculus artigen (CD72) gene	Lycobersion esculentum mRNA for ubiquitin activating enzyme	U.meitanogaster Chc mRNA for clathrih heavy chain	V.alginolyticus sucrase (scrB) gene, complete cds	V. alginolyticus sucrase (scrB) gene, complete cds	AZDDUR.S1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:431799 3	Accounts 1 Soares fetal liver spleen 1NFLS S1 Home saplens cDNA clone IMAGE:431799 31	haitus navegicus initation factor-2 kinase (elF-2a) mRNA, complete cds	N. sabadum Neir-4A13 mRNA	Gallus gallus gene for melanocartin 2-receptor, complete cds	I-OWIDOX Virus, complete genome	Glardia Intestinalis variant-specific surface protein (vsp417-8) gene vsp417-8/A allole	Human mRNA for KIAA0309 gene, partial cds	Homo saplens IA-2 gene, Intron 18	L.mesenteroides gene for sucrose phosphorylase (EC 2.4.1.7)	NUCLEOSIDE TRIPHOSPHATASE I (NUCLEOSIDE TRIPHOSPHATE PHOSPHOHYDROLASE II) ABH	(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	Schaill luberosum cold-stress inducible protein (C17) gene, complete cds	Joyneigus cunculus KING-finger binding protein mRNA, partial cds	AV743773 CB Homo septens cDNA clone CBMAFD06 5'	POZ 18381F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4276381 6	naturs novegicus cytocentrin mRNA, complete cds Dbs=Dbj guanine nucleotide exchange factor homolog fmice. 32D murine hemonolotic cell lico. —BNA assec	מלו שוני שוני שוני שוני שוני שוני שוני שונ
ıgle Exon Pı		Database Source		L N	Z	Z	EST_HUMAN	SWISSPROT	L I	I N	OWISSING!	- L	Į.	112	2 2	114	11-	TOUR TOUR	NAINOL 181		1	i di		LN	Į.	2 12	-	PORGESIMS	TN	L	EST ULIVANI	FRT LIMAN			
เรื่อ	Top Hit Acessian		0.007.0004	7.4E-01 ABUZ1490.2	AB02148	6763217 NT	7.4E-01 AI472641.1	E-01 P09710	E-01 AE001166.1		7.3E-01 135772 1	7.3E-01 L35772 1	7.3E-01 AJ011418 1			T	7.3E-01 AAR78040 4	T	T	X79140 1		7.2E-01 AE108100 1	I^{-}		04 A E408003 4	T		01 23066	-	-	T	T	T		
	Most Similar (Top) Hit	BLAST E Value	7 45 0	1.40	7 45 64	7 july 1	7.41.0	7.37.0	7 35 0	7.38-01	7.3E-01	7.3E-01	7.3E-01	7.3F.01	7.3E-01	7.3F-01	7.3E-04	7.3E-01	7.2E-01	7.2E-01	7.2E-01	7 27.01	100	7.20.07	7 25 04	7.27.01		7.2E-01	7.2E-01	7.2E-01/	7.2E-01	7.2E-01	7.2E-01	7.28.04	
	Expression	Signal	1 65	201	28.5	1000	0 0	0 0	437	101	5.92	5.92	0.67	0.66	7.84	7.84	3.83	3.83	3.89	2.32	1.27	1.38	93.0	3 6	0 7	2.65		0.74	0.88	1.11	0.48	2.33	4.02	1.27	
	ORF SEQ		37582	37589			28413	28838	30018	30373	32278	32277	32735	33118	33210	33211	37361	37362		27398	27920	28473	28851	28894		30083		30410	32864	34183		36100	36568	37049	
	Exon SEQ ID	ë	24260	L	L		L		L	17769	1	1	25103	20040	20121	20121	24056	24056	13583	14685	15181	15829	16201	18354	16785	17450	-	17794	19800	21046	21553	22887	23330	23774	
	Probe SEQ ID	Ö	11665	11665	11900	12008	2888	4575	4652	5040	6511	6511	6894	7359	7445	7445	11407	11407	912	1930	2463	3063	3445	3801	4040	4718		5075	7112	I	- 1	10239	10639	11104	

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			ase channel (ryanodine receptor) alpha		exons 15-16			4296344 5	4296344 5	gene, complete cds	clone IMAGE:202961 3'			3898495 5		3E:731109 3'			cDNA clone IMAGE:288708 3' similar to	cDNA clone IMAGE:288708 3' similar to		97	genome	system (PTS) system, mtlA, mtlR, mtlF,		system (PTS) system, mtlA, mtlR, mtlF,					cds and translational regulator gene, partial
	Top Hit Descriptor	Aeropyrum pernix genamic DNA, section 6/7	Rana catesbelana mRNA for builfrog skeletal muscle calcium release channel (ryanodine receptor) alpha	isoform(RyR1), complete cds	Homo sapiens partial TCF-4 gene for T-cell transcription factor-4, exons 15-16	Mus musculus otogelin (Otog), mRNA	Mus musculus otogelin (Otog), mRNA	602155438F1 NIH_MGC_83 Homo septens cDNA clone IMAGE:4296344 5	602155438F1 NIH MGC 83 Homo sepiens cDNA clone IMAGE:4296344 5	Drosophila melanogaster 6-pyruvoyltetrahydropterin synthase (pr) gene, complete cds	yq89d09.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:202961 3	RC1-BT0567-301299-011-d09 BT0567 Homo saplens cDNA	RC1-BT0567-301299-011-d09 BT0567 Homo saplens cDNA	601496330F1 NIH MGC 70 Homo sapiens cDNA done IMAGE:3898495 5	Human T-cell receptor germline gamma-chain J2 gene	Zu06h11.s1 Soares testis NHT Homo sapiens cDNA clone IMAGE 731109 3'	Homo sapiens mRNA for KIAA0614 protein, partial cds	Homo seplens mRNA for KIAA0614 protein, partial cds	yz73e07.s1 Soares, muttple_solerosis_ZNbHMSP Homo sepiens cDNA cione IMAGE.288708 3' similar to contains Alu repetitive element;	yz/3e07.s1 Soares_muttple_sdercs!s_2NbHMSP Homo sapiens cDNA clone IMAGE:288708 3' similar to	Homo saniens chromosome 21 seament HS21C101	Arabidopsis thaliana mRNA for chlorophyll b synthase complete cds	Escherichia coli K-12 MG1855 section 143 of 400 of the complete genome	Clostridium acetobutylicum mannitol-specific phosphotransferase system (PTS) system, mild, mills, mills, mills,	and mtD genes, complete cds	Clostridium acetobutylicum mannitol-specific phosphotransferase system (PTS) system, mtlA, mtlR, mtlF,	and mtiD genes, complete cds	Danio rerto complement factor B mRNA, complete cds	AV763842 MDS Homo saplens cDNA clone MDSCHE04 5"	AV763842 MDS Homo sapiens cDNA clone MDSCHE04 5'	Candida albicans squalene epoxidase (CAERG1) gene, complete cds and translational regulator gene, partial cds
	Top Hit Database Source	N		Ę	IN	Ĭ	¥	EST HUMAN	EST_HUMAN	L'N	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	EST HUMAN	LN	FN	EST_HUMAN	TOTAL TOTAL	LN FIN	LZ	LN PN		NT		NT	LN	EST_HUMAN	EST_HUMAN	NT
5	Top Hit Acession No.	7.2E-01 AP000063.1		21 D21070.1	7.1E-01 AJ270777.1	7305360 NT	7305360 NT	1 BF681034.1	1 BF681034.1	1 U36232.1	1 H54244.1	1 BE074185.1	BE074185.1	BE904405.1	M12961.1	7.1E-01 AA421492.1	7.0E-01 AB014514.1	7.0E-01 AB014514.1	7.0E-01 N62412.1	JR2412 1	2								7.0E-01 AV763842.1	7.0E-01 AV763842.1	
	Most Similar (Top) Hit BLAST E Value	7.2E-01		7.1E-01	7.1E-01	7.1E-01	7.1E-01	7.1E-01	7.1E-01	7.1E-01	7.1E-01	7.1E-01	-	1 —	7.1E-01	7.1E-01	7.0E-01	7.0E-01	7.0E-01	7 0E-04 N82412 1	7.0F-01	7.0E-01	7.0E-01		7.0E-01 U53868.1		7.0E-01 U53868.1	7.0E-01 U34662.1	7.0E-01	7.0E-01	6.9E-01 U69674.1
	Expression Signal	2.9		12.73	11.76	3.18	3.18	1.63	1.63	7.68	0.56	0.93	0.93	1.43	1.22	2.21	0.99	66.0	1.13	1.13	1 78	1.11	11.76		0.57		0.57	0.49	1.94	1.94	11.02
	ORF SEQ ID NO:					29655		31585		32580	33916	34469			36172		26824	26625	27907	S/DUZ/C			_		35064			36400	37006	37007	26380
	SEQ ID NO:	24798	L.		[18645	19550						25205	13958	13958	15169	15169	17719	1	20970	1	21895		- 1	23172	23734	23734	13715
	Probe SEQ ID NO:	12422		9/9	3059	4184	4184	2858	5858	. 6850	8091	9635	8635	9755	10309	12211	1207	1207	2450	2450	4998	5862	8276		9216		9216	10526	11064	11064	949

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Top Hit Descriptor	Candida albicans squalene epoxidase (CAERG1) gene, complete ods and translational regulator gene, partial	DD28a09 st NCI COAB CLAIL	Chlemylle mindering conf. 2 for the control of the	Smentyde innederin, section 3 of 85 of the complete genome	Prencinosiona beroneri BbNA3 mRNA for notochord actin, complete cds	S01177733251 AILU 1905 27 III	Strong Asset I Nin _ WGC_17 Home sapiens cDNA clone IMAGE:3632328 6	Archidansis thelians DNA -	reconstruction and the control of th	A abloopsis maliana DNA chromosome 4, contig fragment No. 69	Enternospa dispar cation transporting ATPase (atpase) gene, partial cds	muse acuminata pectate Iyase 1 (PL1) mRNA, complete cds	Musa acuminata pectate Iyase 1 (PL1) mRNA, complete cds	Homo sapiens DAN gene, complete cds	Homo sepiens DAN gene, complete cds	FORKHEAD BOX PROTEIN C2 (FORKHEAD-RELATED PROTEIN FKHL14) (MESENCHYME FORK	Sindia In (With-1 PROIEIN) (TRANSCRIPTION FACTOR FKH-14)	Streetheamelis carbamate Khase gene, complete eds	Syndronous sp.: P.C.Costus complete genome, 27/27, 3418852-3573470 3/75e06.s1 Soares, parathyroid tumor NDHPA Homoseniens, PNA, Alexania, 114007-4, 200552	35:X56411 ma1 ALCOHOL DEHYDROGENASE CLASS II PI CHAIN (HUMAN)	તથ(hooded) prolactin gene : exon ili and flanks	domo saplens mRNA for KIAA1345 protein, partial cds	Stagonospora avenae bgf1 gene for beta-glucosidase. expns 1.4	Nagonospora avenae bgl1 gene for beta-quicosidasa exeme 1.4	Aus musculus zinc finger protein (Pens) mRNA complete	flue musculus zino finger protein (Peas) mRNA complete cas	Aus musculus major histocompatibility complex region NG27, NG28, RPS28, NADH oxidoreductase, NG29, IFC1, Fas-binding protein, BING1, tapasin, RaiGDS-like, KE2, BING4, beta 1,3-galactosy transferase, and	Mus musculus major histocompatibility complex region NG27 NG28 DBS29 NATE:	KIFC1, Fas-binding protein, BING1, tapasin, RalGDS-like, KE2, BING4, beta 1,3-galactosy transferase, and RPS18 genes, complete cds: Sarm21 name mouth.
Top Hit Database Source	Ę			IN	Į.	EST HIMAN	LN LN	Į	Į.	EV.	1	1	1						T	T_HUMAN									
	U69674.1	AA583530.1	AE002271.2	AB035662 1	Y18278 1	-	T	Τ	F		T		T				T				T								6.8E-01 AF110520.1
Most Similar (Top) Hit BLAST E Value	6.9E-01	6.9E-01	6.9E-01	6.95-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.9E-01	6.95-01	6 9F-01	2010	6.9E-01		6.8E-01/	6.8E-01		0.87-01	0.00-01	0.00-01	0.05-01	6.8E-01	6.8E-01 A	6.8E-01 A	6.8E-01		6.8E-01
Expression Signal	11.02	2.74	1.97	0.91	0.82	1.36	0.65	2.96	2.96	0.79	0.59	0.69	2.38	0000	8,30	3.01	1.05	66.0	,	2 4	2 4	5 6	76.	1.92	2.4	2.4	1.38		1.36
ORF SEQ ID NO:									33701		35431	35432	37172	37173	21.5		26369		27046	28901	35375	36962	20000	20000	3/000	3/001	37527		37528
	1		\perp			19050	20360	20574	20574	21758	22247	22247	23886	23886		26197	13704	15389	14358	17268	22191	23690	22600	207700	97/59	63/63	24205		24206
SEO DO NO:	949	1287	3213	5694	2900	6277	7697	7879	7879	8066	9594	9594	11223	11223		11878	937	2880	2832	4533	9538	11027	11027	44080	1406	2	11607		11607
	Exon ORF SEQ Expression (Top) Hit Top Hit Acession No. Signal BLASTE No. Source	No: CAF SEQ Expression (Top) Hit Top Hit Acession Top Hit	Signal Apple Signal Apple Ap	Exon Nos: Signal No: 13715 ORF SEQ ID No: Signal 11.02 Expension (Top) Hit Top Hit Acession No: Signal No: Signal No: Source Native No: Source No	Exon Nos: Signal No: Signal Signal Most Similar Top Hit Acession No: Top Hit Acession Database Source Top Hit Acession No: Top Hit Acession Signal Top Hit Acession No: Top Hit Acession No:	Exon Nos: Signal No: No: No: Source No: No: No: No: No: No: No: No: No: No:	Exon Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Signal Nos: Nos: Signal Nos: Nos: Nos: Nos: Nos: Nos: Nos: Nos:	Exon Nos: Signal No: Signal Signal Most Similar Top Hit Acession No: Top Hit Acession No: Top Hit Acession Signal Top Hit Acession No: Top Hit Acession No:	Exon Nos: Signal No: Signal Signal 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Interval NT 23886 37172 2.38 6.9E-01 Ace0319.1 Interval NT Interval NT<	Exon ORF SEQ Expression Most Similar Top Hit Acession Aces Aces Aces Aces Aces Aces Aces Aces	Exon ORF SEQ Expression Most Similar Top Hit Acession Aces Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acession Ace Hit Acess	Exon Nost Signal ORF SEQ Nossion Nost Similar Top Hit Acession Nost Similar Top Hit Acession Database Top Hit Top Hit Acession Source 13715 26381 11.02 6.9E-01 Usge74.1 NT 14037 26708 2.74 6.9E-01 Usge74.1 NT 14037 26708 2.74 6.9E-01 Ace0s271.2 NT 14037 26708 2.74 6.9E-01 Ace0s271.2 NT 18488 31409 0.91 6.9E-01 Ace0s271.2 NT 19650 32027 1.36 6.9E-01 Ace0s271.2 NT NT 20574 33701 2.9E-01 Ace0s271.2 NT NT NT 20574 33701 2.9E-01 Ace0s31.3 NT NT NT 22247 33701 2.9E-01 Ace0s19.1 NT NT NT 23886 31772 2.3B-0.6E-01 Ace0s19.1 NT NT NT 23886 37172 2.3B-0.6E-01 Deg013.1 NT NT H 23886 37173 2.3B-0.6E-01 Deg013.1 NT	Exon Nost Signal ORF SEQ Notes signal Signal (Top) Hit Top Hit Acession No. Top Hit Acession Source Sou	Exon Nost Signal ORF SEQ Notes signal Signal Notes Signal Most Similar Top Hit Acession <	Exon Nost Signal ORF SEQ Notession of Crop Hit Top Hit Acession of Signal of	Exon Nost Signal ORF SEQ Notes signal Signal (Top) Hit Top Hit Acession No. Top Hit Top Hit Acession Source Nation Top Hit Top Hit Acession Source So	Expn No: CRF SEQ Seg ID No: Expression Signal (Top) Hit Top Hit Acession Top Hit Acession Politics Top Hit Acession Politics Top Hit Acession Politics Top Hit Acession Source Top Hit Acession Source	Expn No: NO: NO: NO: NO: NO: NO: NO: NO: NO: NO	SECO DRF SEQ Expression Top Hit Acession SEQ ID ID NO: Signal Crop Hit Acession Top Hit Acession Top Hit Acession Top Hit Acession Database	Expn NO: CRF SEQ Seques Expression Signal Most Similar Fraid Top Hit Acession Pulse Top Hit Acession Pulse Top 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Organia I.1.03 S.BC.O. Organia I.1	SEC D ORF SEQ Signal Most Similar Top Hit Acession Database No. 13715 26381 11.02 6.9E-01 Accession Database Source Source Source Accession Co.	
Page 41 of 536 Table 4 Single Exon Probes Expressed in Brain

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Table 4
Single Exon Probes Fxnmssed i

Single Exon Probes Expressed in Brain	Most Similar (Top) Hit Acession (Top) Hit Acession (Top) Hit Documents (Top) Hit Document	26020 18.23 6.5E-01 M75140 1	28833 4.26 8.5F-01 ARA41228 1 NT	29815 4.23 6.5E-0.1 A 127228E 4 NIT	29843 0.78 6.5F-04 A1 484590 2 N.T.	N Template	7726 30329 2.6 6.5E-01 028921.1 NT Phaseolus vulgaris ATPase camma subunit mRNA Arrichae and Arr	30843 1.77 6.5E-01 P18480 SWISSEDOT	31337 0.62 6.5E-01 AL 163249 7 NT	32400 1.5 6.5E-01 DAR348 1 NIT	33340 0.84 6.5E-01/A/799882-1 FINANI	0.8 6.5E-01 T78904.1 EST LIMAN	36094 1.96 6.5E-01 AF119878 1 NIT	36460 2.68 6.5E-01 H87583.1 EST HUMAN	36518 3.5 6.5E-01 AA601287.1	3.93 6.5E-01/AU138078.1 FET HIMAN	375/A 22 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2.07 A SE-01 AF014115.1	1.81 6.5E-01 272448 4 NIT	25694 8.05 6.4E-01 U48848.1 NT	28043 1.16 6.4E-01 AF1611841 NT	28855 2.16 6.4E-01 U48854.2 NT	29230 1.08 6.4E-01 AB046827.1 NT	34347 1.82 6.4E-01 AE001247.1 INT	35848 8.6 6.4E-01 U82828.1 NT	35864 1.22 6.4E-01 BF670405.1 EST HUMAN	5.99 6.4E-01 AV759212.1 EST HIMAN	25858 4.58 6.3E-01 P05228 SWISSPROT	26938 2.26 8.3E-01 U32689.1 NT	27623 2.02 6.3E-01 U81136.1	28035 3.51 6.3E-01 U75331.1 NT	28036 3.51 6.3E-01 U75331.1	
		26020	28833	29615	29643	-	30329	30843	31337	32400	33340	 -	36094	36460	36518		37518			25694	28043	28855	29230	34347	35848	35864		25858	26938	27623	28035	28036	
	Exan SEQ ID NO:	13388	16183	16990	L	L	17728	25087		. 1	,		22881	23228	23280	23381	24198	24710	25146	13054	15307	16205	16593	21202	22637	22652	24777	13211	13308	14889	16297	15297	
	Probe SEQ (D NO:	910	3428	4249	4277		2003	5367	5627	6825	7366	9737	10233	10529	10685	106901	11599	12267	12504	245	2683	3449	3842	8510	8866	1000	12382	425	522	2159	2683	2583	

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5973	18755	31716	0.94	6.3E-01	1 BE093906.1	EST_HUMAN	PM0-BT0757-010500-002-a05 BT0757 Homo sapiens cDNA
6504	19269	32271		6.3E-01	L27798.1	NT	Streptococcus dysgalactiae (mag) gene, complete cds
6504	19269	32272		6.3E-01	L27798.1	LN	Streptococcus dysgalactiae (mag) gene, complete cds
8419				6.3E-01	BE902044.1	EST_HUMAN	601676889F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3959351 5'
8784		34624	96'0		S62927.1	LZ LZ	glycoprotein Illa (Alu 1 and 3 fusion junction) [human, Genomic Mutant, 300 nt]
9120		34975			BF216984.1	EST_HUMAN	601884050F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4102596 5'
9320		35159		6.3E-01	9627521 NT	Z	Varida virus, complete genome
9320		35160			9627521 NT	Z Z	Varida virus, complete genome
9838			19.0	6.3E-0	11 AE002329.2	N	Chlamydia muridarum, section 59 of 85 of the complete genome
10326	22973	36193			6.3E-01 Z73003.1	ĮN.	S.cerevisiae chromosome VII reading frame ORF YGR218w
10427		36294	1.19	6.3E-01	AE000313.1	N	Escherichia coli K-12 MG1655 section 203 of 400 of the complete genome
10456			0.45	6.3E-01	6.3E-01 AW795395.1	EST_HUMAN	PMC-UM0018-130500-003-g12 UM0018 Homo sapiens cDNA
							nr09h06.s1 NCI_CGAP_Co10 Homo sapiens cDNA clone IMAGE:1161371 3' similar to TR:002916 002916
10993		36924		6.3E-01	6.3E-01 AA877715.1	EST_HUMAN	HLARK.;
11308		37268		6.3E-01	6.3E-01 Al904160.1	EST_HUMAN	CM-BT043-090299-046 BT043 Homo saplens cDNA
11402	24051	37355		6.3E-01 P47003	P47003	SWISSPROT	HYPOTHETICAL 13.7 KD PROTEIN IN INO1-IDS2 INTERGENIC REGION
11581		37495		6.3E-01 P36073	P36073	SWISSPROT	HYPOTHETICAL 15.3 KD PROTEIN IN VMA12-APN1 INTERGENIC REGION
11988		30607	4.37	6.3E-01	9910293 NT	NT	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA
12078	_		1.45	6.3E-01	6.3E-01 AF105227.1	IN	Homo saplens 3'-phosphoadenosine 5'-phosphosulfate synthetase (PAPSS) mRNA, complete cds
12283	Ì.		2.93	6.3E-01		TN	C.limicola pscD gene
5780		31499	2.31	6.2E-01		SWISSPROT	HYPOTHETICAL 142.5 KD PROTEIN C23E2.02 IN CHROMOSOME I
7394	20073		3.44	6.2E-01		IN	Mus musculus calclum-sensing receptor related protein 4 (Casr-rs4) mRNA, partial cds
							Mus musculus chromosome X contigA; putative Magaa9 gene, Caltractin, NAD(P) steroid dehydrogenase
7443	25114	33209	1.33	6.2E-01		Z	and Zinc tinger protein 185
8200	- 1	34031		6.2E-01	6.2E-01 H72255.1	EST_HUMAN	ys01e08.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:2135423'
							Lycopersioon esculentum cytosolic Cu,Zn supercodde dismutase (Sod) gene, partial cds; and dehydroquinate
8755	- 1	34595		6.2E-01	1	NT	dehydratase/shikimate:NADP oxidoreductase gene, complete cds
8349	1	33540		6.2E-01	BE562687.1	EST HUMAN	601336146F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3690010 5'
9410			2.55	6.2E-01	6.2E-01 M24461.1	NT	Human pulmonary surfactant-associated protein SP-B (SFTP3) mRNA, complete cds
9978	22626	35834	6.2	6.2E-01		NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 23
10121	22769	35982	0.5	6.2E-01	11420793 NT	Į.	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1), mRNA
10121	22769	35983	0.5	6.2E-01	11420793 NT	N	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1), mRNA
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| Тор Hit Descriptor | NON-STRUCTURAL POLYPROTEIN [CONTAINS: RNA-DIRECTED RNA POLYMERASE; THIOL PROTEASE P3C; HELICASE (2C LIKE PROTEIN) | NON-STRUCTURAL POLYPROTEIN (CONTAINS: RNA-DIRECTED RNA POLYMERASE; THIOL PROTEASE P3C; HELICASE (2011 IKE PROTEIN): COAT PROTEIN! | Mus musculus secreted acidic cystene rich alvoporotein (Sparc) mRNA | Caenorhabditis elegans N2 CeMvoD (hih-1) alternatively snilload papes, complete cde | Rat TRPM-2 gene, complete cds | Rat TRPM-2 gene, complete cds | xd50h03.x1 NCI_CGAP_Ov23 Homo septens cDNA clone IMAGE:2597237 3' sImilar to gb:X12671_rna1
HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 (HI IMAN): | SUSHI REPEAT-CONTAINING PROTEIN SRPX PRECURSOR (DRS PROTEIN) (DOWN-REGULATE
BY V-SRC) | Arabidoosis thaliana outsitive zinc transporter (ZIP4) mRNA complete and | Homo sabiens mitogen-activated protein kinasa kinas | Homo sapiens mitoden-activated protein kinasa
kinasa kinas | Homo saplens G-protein coupled receptor EDG-7 mRNA, complete cds | Homo sapiens G-protein coupled receptor EDG-7 mRNA complete cds | Pseudomonas aerudinosa PA01, section 13 of 529 of the complete general

 | Homo sapiens dopamine transporter (SLC8A3) gene, complete orts | hysturonan-binding protein-thepstocyte growth factor activator homolog fluman, plasma, mRNA, 2408, m | hvaluronar-bindina protein=thematocode arough factor activator homolog formula | Mus musculus Codas mRNA for tone IV collades aches schein complete such | M.mazel orfA, orfB, and orfC of archaeal ABC-transporter system | Homo saplens DNA for amyloid precursor
protein, complete ods | Homo sapiens adaptor-related protein complex 3 mi 2 subunit (Cl A2n) mRNA
 | Human respiratory syncytial virus strain CH33-53b attachment protein (G) gene complete cde | Viral hemorrhagic septicemia virus N. P. M. G. Nv. L. genes. French strain 07-71 | Homo sapiens Notch3 (NOTCH3) gene, exams 28, 27, and 28 | D(2) DOPAMINE RECEPTOR | UI-H-BI1-aeb-a-10-0-U.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718619 3'
 | Musca domestica insecticide-susceptible strain voltage-sensitive sodium channel mRNA, complete ods |
| Top Hit
Database
Source | SWISSPROT | SWISSPROT | NT | PN | LZ
LZ | L | EST_HUMAN | SWISSPROT | Ę | Į. | Z L
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| Top Hit Acession
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 | AF236117.1 | Γ |

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| Most Similar
(Top) Hit
BLAST E
Value | 6.2E-01 | 6.25-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01
 | 6.1E-01 | 6.1E-01 | 6.1E-01

 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.1E-01 | 6.0E-01
 | 8.0E-01
 | 6.0E-01 | 6.0E-01 | 6.0E-01 | 6.0E-01 | 8.0E-01
 | 6.0E-01 U38813.1 |
| Expression
Signal | 5.2 | 5.2 | 4.38 | 1.15 | 4.02 | 4.02 | 0.64 | 0.72 | 3.27 | 1.09 | 1.09
 | 18.74 | 18.74 | 0.93

 | 1.06 | 2.57 | 2.57 | 2.28 | 1.57 | 1.24
 | 3.09
 | 1.91 | 0.92 | 1.09 | 2 | 2.86
 | 2.78 |
| ORF SEQ
ID NO: | 36305 | 36306 | | | | | 32702 | 32751 | 33962 | 34528 | 34529
 | 35153 | 35154 | 35597

 | 35797 | 37855 | 37656 | 30899 | | 25903
 | _
 | 28785 | 29180 | | 30628 | 30839
 | 32210 |
| Exan
SEQ ID
NO: | 23080 | 23080 | 15114 | 18248 | 19514 | 19514 | 19656 | 19897 | 20826 | 21386 | 21386
 | 21982 | 21982 | 22393

 | 22594 | 24331 | 24331 | 25159 | 24977 | 13267
 | 13331
 | 14089 | 16547 | 16905 | 18007 | 18166
 | 19213 |
| Probe
SEQ ID
NO: | 10434 | 10434 | 2393 | 5449 | 6770 | 6770 | 6920 | 7005 | 8132 | 8694 | 8694
 | 9315 | 9315 | 9742

 | 9946 | 11738 | 11738 | 12074 | 12694 | 482
 | 648
 | 1341 | 3795 | 4165 | 5189 | 5353
 | 6445 |
| | SEQ ID ORF SEQ Signal (Top) Hit Top Hit Acession (Top) Hit Top Hit Acession (Top) Hit Top Hit Acession Database No. | Expn
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Source 23080 36306 6.2 6.2E-01 P27410 SWISSPROT
SWISSPROT 15114 4.38 6.1E-01 P27410 SWISSPROT
SWISSPROT | Expn
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SWISSPROT 15114 4.38 6.1E-01 P27410 SWISSPROT
SWISSPROT 11514 4.38 6.1E-01 M59940.1 NT | Expn
SEQ ID
NO: ORF SEQ
Signal Expression
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BLAST E
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NO: Top Hit Accssion
NO: Top Hit Accssion
Source 23080 36306 6.2 E-01 P27410 SWISSPROT 15114 4.38 6.1 E-01 P27410 SWISSPROT 18248 31137 1.15 6.1 E-01 MT 19514 32540 4.02 6.1 E-01 M694733.1 NT | Expn
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Page 45 of 536 Table 4 Single Exon Probes Expressed in Brain

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тор Hit Descriptor
6583			0.68		Q04912	SWISSPROT	MACROPHAGE-STIMULATING PROTEIN RECEPTOR PRECURSOR (MSP RECEPTOR) (P185-RON) (CDW 138) (CD136 ANTIGEN)
7254				6.0E-01	AJ277661.1	N.T.	Homo sapiens partial LMO1 gene for LIM domain only 1 protein, exon 1
8023			66.4	6.0E-01	P02835	SWISSPROT	SEGMENTATION PROTEIN FUSHI TARAZU
8023	20718	33851	4.39	6.0E-01	P02835	SWISSPROT	SEGMENTATION PROTEIN FUSHI TARAZU
9723	22374	35574	1.61	10-30:9	AB008193.1	5	Homo sanjans namae for ferikritiana R4 recentive B1 T2 I salizations B4 recentive B4
10173			1.48		Q01497	SWISSPROT	PEROXISOMAL MEMBRANE PROTEIN PERB (PEROXIN.3)
10990	Ш		1.49		AJ131892.1	Ę	Gallus gallus mRNA for Hyperion protein, 419 kD isoform
10990			1.49	6.0E-01	AJ131892.1	Ę	Gallus gallus mRNA for Hyperion protein, 419 kD Isoform
11540		37449	3.77		AI420623.1	EST_HUMAN	#08f07.xt NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE: 2095621.31
12354		31060	2.25		11421663		Homo saplens nuclear factor (enythroid-derived 21-like 3 (NFE2I.3), mRNA
12455	ı		2.6	6.0E-01	AA706087.1	EST HUMAN	2998905.s1 Soares fetal liver spleen 1NFLS S1 Home sablens cDNA clone IMAGF 462776 3
12639		30815	3.04	6.0E-01	B055303 NT	٧.	Mus musculus cGMP-inhibited phosphodiesterase (Pde3a), mRNA
12684			2.08		BE157617.1	EST_HUMAN	RC1-HT0375-030500-015-c03 HT0375 Homo sapiens cDNA
88			1.38		U32701.1	IN	Hæmophilus influenzae Rd section 16 of 163 of the complete genome
3264	I		2.29		AL163267.2	ŢN	Homo sapiens chromosome 21 segment HS21 C067
3264	_	28676	2.29	5.9E-01	AL163267.2	NT	Homo saplens chromosome 21 segment HS210087
4196	16937		4.21		AF162756.1	NT	Rattus norvegicus cenexin 2 mRNA, partial cds
6373	19142		1.55		AF065440.2	-N	Homo saplens low density lipoprotein receptor-related protein II (I RP2) nems even 1 and northal sets
7166	19853	32922	1.32	5.95-01	AB023486.1	L	Homo saplens gene for histamine H2 receptor, pramoter recion and complete ods
7296	19979		0.61		X68801.1	ĽN	G.galius gene for skeletal alpha-actinin, exon EF2
7898	20593	33725	0.46		D90911.1		Synechocystis sp. PCC6803 camplete gename, 13/27, 1576593-1719643
8536	21228	34370	0.48		D12922.1	LT	Legionella pneumophila gene for iron superoxide dismutase, complete cds
0443	20100	96300	6				
8	73/67	2000	0.02	3.3E-01	7,	7	Crimingia ucanometrs strain K/UW 31/CX major outer membrane protein (omp1) gene, complete cds
2000	70,77		0.74	5.8E-U1		٦	E6 PROLEIN
10088	22736	35951	1.15	5.9E-01			VASCULAR ENDOTHELIAL-CADHERIN PRECURSOR (VE-CADHERIN) (CADHERIN-5)
10569	23284	36502	2.5			SWISSPROT	THYMIDYLATE KINASE (DTMP KINASE)
10578	23271	36507	1.72		AF197944.1	NT	Xenopus laevis receptor protein tyrosine phosphatase delta (XPTP-D) mRNA, complete cds
10881	23561	36808	2.91			T_HUMAN	PM1-DT0041-190100-002-h03 DT0041 Hamo sapiens cDNA
11149	23816	37099	1.95		28.1		Mus spretus strain SPRET/EI CD48 antigen (Cd48) gene, partial cds
11458	24062	37368	1.58	5.9E-01	P47135	SWISSPROT	JSN1 PROTEIN

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				- 1		1	PYRROLINE-S-CARBOXY/ATE REDUCTASE (P5CR) (P5C REDUCTASE)
7664	20328	33438	2.14	5.7E-01		ISSEROI	Men Territor Kong I from Mach 2 Tapa-1. Isset and Tssch genes, alternative transcripts
7870	20565		0.5	5.7E-01		Т	Mus musculus North I. E. P. C. M. M. Inram Home sapiens cDNA
8279	1		0.47	5.7E-01		HOMAN	Thousand Training Town the Promosome 4 control fragment No. 32
9699	ı	35544	1.19	5.7E-01			Arabidopsis integrated by Arabidose A configuration No. 32
0000	22350		1.19	5.7E-01		\neg	Arabidopsis intalina Livin singmooning transportations (AMACE: 4088810 5)
10475			0.72	5.7E-01		П	60206/712F1 NIT MIGG 20 19110 september 201NA
44083	L		1.52	5.7E-01	01 BE715051.1	HUMAN	MR3-H10/36-180/UU-005-EUZ-II O SOO III O SOO I
1265	1		3.01	5.7E-01	01 BE959722.2	HUMAN	601654814K1 NIH MIGC 37 TIOITIO Septemb COMP. Section 11.
	L	02770	13	5.6E-01	01 AB018283.2	N	Homo sapiens mKNA for Nivavi 40 protein, para e e
230				5.6F-01		NT	Homo sapiens mRNA for KIAA0740 protein, partial cds
ĝ			ľ			Į.	Arabidopsis thaliana DNA chromosome 4, contig fregment No. 13
386				30.0	Ī	1	Chicken TBP gene, exon8, complete cds
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	10000			5.6E-01	-01 AV684703.1	EST_HUMAN	AV684/03 GRC home septents conviction and all
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11997	7 24535			30.0		1	Arabidousis thallana DNA chromosome 4, contig fragment No. 13
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12379	9 24776	9	2.7			SWISSING	ACCIONAGE NIM MGC 81 Homo saplens cDNA clone IMAGE:4271334 5
12773	73 25027	7	4.26		5.6E-01 BF573829.1	ESI HOMAIN	Control of the Control Control of Carbondase beta polypeptide (Poob), mRNA
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	1		80		5 5E-01 P03341	SWISSPROT	PROTEIN P30; NUCLEOPROTEIN P10]
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Single Exon Probes Expressed in Brata	Top Hit Descriptor		EST02835 Fetal brain, Stratagene (cat#938206) Homo saplens CINA class LIEBCO37	601811077R1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4054003 3'	Homo saplens KIAA0929 protein Msx2 Interacting nuclear target (MINT) hamalog (KIAA0929)	Homo sapiens KIAA0928 protein Mev2 interesting	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE). HrpW (hrpW), and GstA (gstA) genes.	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE). Hrnw (hrnw).	complete cds; and unknown genes	CV 4-INNUG40-070400-160-c04 NN0040 Homo sapiens cDNA	Onscribilia pneumoniae AR39, section 74 of 94 of the complete genome	Cocypinal meanogaster mRNA for 15,15' beta carotene dioxygenase (beta-diox gene)	DAYS CALLOS OLD glucuronosyltransferase gene, complete cds	Rathis possession	remains rely algebra gate for 11S11, complete cds	S. cerevisiae RIR3 name encoding PDP	Cerentsiae RIR3 and control in the C	MITOCHONDRIAL TRIFUNCTIONAL ENZYME ALPHA SUBUNIT PRECURSOR (TP-ALPHA) [INCLUDES: LONG-CHAIN ENOYL-COA HYDRATASE - I ONG CHAIN 9 LYCHOOLOGO	DEHYDROGENASE]	6UZ076545F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4243890 F	NI KATE REDUCTASE [NADPH] (NR)	LAMININ ALPHA-2 CHAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN)	WISTOOK IN CARAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN) WISTOOK IN CLEAP LUTH HOMO SEPIENS CONA CIONA IMAGE: 2427426 CH. 1	IUMAN);	hydroxylase (CYP21B), complement component C4 (CAB) G11 helicaes (SYICH) CA	(Bf), and complement component C2 (C2) genes,>	Brassica cleracea var. capitata phospholipase D2 (PLD2) gene, complete cds	Brassica oferacea var. capitata phospholipase D2 (PLD2) gene, complete cds
igle Exon Pro	Top Hit Database Source	100	ESI HUMAN	EST_HOMAN	MT	Ä	'n	1	T LI MAAN	Т			┰	L	Т	NO.	E		SWISSPROI	T	7	1	T	TO NOWAN				
<u> </u>	Top Hit Acession No.	-04 T05047 4	04 BE420507 4	Dr 128007.1	7657266 NT	7657266 NT	01 AF232006.1	5.4E-01 AF232008 1		T	Γ		L		l		Z21619.1	084478	, 0	T					A E010442 4	Ī	AF113010 1	1632B
	Most Similar (Top) Hit BLAST E Value	5.5F-01	5 5E-04		6.4E-01	5.4E-01	5.4E-01	5.4E-01	5.4E-01/	5.4E-01/	5.4E-01	6.4E-01	5.4E-01	5.4E-01 A	5.4E-01 B	5.4E-01 Z21619.1	5.4E-01 Z	74 F	5.4F-01 RI		5.4E-01.0	5.4E-01.06	5 4F_01 A II	_	5.35-04	_	6.3E-01 AF	6.3E-01
	Expression Signal	0.96	1.65	3	18,4	4.91	1.16	1.16	3.41	3.43	1.91	0.92	0.74	0.83	0.87	0.81	0.81	.48	2.09	2.87	3.08	3.08	3.5		1.54	1.01	1.01	6.83
	ORF SEQ ID NO:	L	37033			25598	26980	25981	26664		27719	30402	31278	31845	32710	32893	35834	32997	-	36948	37541	37542		-	26921	27596	27597	28230
	SEQ (D		23757	12955	L	12955	13352	13352	13997	14830	14980	17786	18368	18876	900	19920	200	19922	22540	23687	24218	24218	24499	_	13287	14866	14866	15491
	Probe SEQ ID NO:	10276	11087	140	;	140	571	571	1248	2089	2262	8	8	860	0760	23 128		7237	88	11015	11621	1621	11944				2136	-1

Page 49 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	Homo saplens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA	Lown carlens secreted C-type lectin precursor (LSLCL) gene, complete cds	Myroplasma denitalium section 9 of 51 of the complete genome	Address overy tumor NbHOT Homo sapiens cDNA clone (MAGE:740711 5	2474.11.2.)? Common North Home Septems cDNA clone IMAGE:740711 5	7.42nn9 r1 Spares NhHMPu S1 Homo sepiens cDNA clone IMAGE:686112 5	2142209 1 Spares NhHMPu S1 Homo sepiens cDNA clone IMAGE:666112 5	7-72-12 v. NCI CGAP Pr28 Home sapiens cDNA clone IMAGE:3288118 3' similar to gb:J02783	PROTEIN DISULFIDE ISOMERASE PRECURSOR (HUMAN); PROTEIN DISULFIDE Date United For Services (NA Clone IMAGE:3288118:3' similar to gb:J02783	PROTEIN DISULFIDE ISOMERASE PRECURSOR (HUMAN):	Roridula gorganias ribulose 1,3-bispriospriate da booynood (15-5) 5-1-1 f	Crind-pract process. 7271c12.x7 NCI CGAP_Lu24 Homo sapiens cDNA clone IMAGE: 3' similar to contains element MER29	repetitive element;	repetitive element;	wagabozyt ncj_cgar_mens hame sapens contractions are the polymenter of precursor; sw.coxa human p20674 cytochrome c oxidase polymenter.	601339867F1 NIH MGC_53 Homo sapiens cDNA clone IMAGE:3682168 5	RASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN	PRECURSOR (HSPG) (PERLECAN) (PLC)	APOLITIOPROTEIN D PRECURSOR (HUMAN); APOLITIOPROTEIN D PRECURSOR (HUMAN); APOLITION OF THE PROTEIN THE	DIOSOPHIBA HIGHAN OF ACTIVATED T CELLS 5 (T CELL TRANSCRIPTION FACTOR NFATS) (NF-ATS)	(REL DOMAIN-CONTAINING TRANSCRIPTION FACTOR NFATS)	Homo saplens phospholipid scrampiase 1 gene, continued out	Homo sapiens curonosario et regimento de la constanta de la co	Homo sapiens minya lid Nazyot of promiting the property of property of the pro	Ontarrydophilia abortos organicas complete cds	Robusta vincera strain T4 cDNA library under conditions of nitrogen deprivation	am77g05.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:1616504 3	
Dani I IIOVI BIBINO	Top Hit Database Source				Т	HOMAN	Т	7	EST HOWAN	EST_HUMAN	EST HUMAN			EST_HUMAN	EST_HUMAN	EST HIMAN	Т	Т	SWISSPROT	EST_HUMAN	2	SWISSPROT	LN L	NT	LN	LN.	LN L	EST HUMAN	
2	Top Hit Acession No.	TIA	4506320	-				5.3E-01 AA193672.1	01 AA193672.1	.01 BE645620.1	.01 BE645620.1		-01 L01950.2	5.3E-01 BF433956.1	-01 BF433956.1	A 105 4040 4	5.3E-01 A1934Z10.1	BE566291.1	5.3E-01 Q05793	5.3E-01 AA916053.1	5.2E-01 L20770.1	5 2F-01 Q9WV30	5.2E-01 AF224492.1	AL163285.2	5.2E-01 AB018283.2	5.2E-01 U65942.1	E-01 D73443.1	5.2E-01 AL116780.1	AA304 103. I
-	Most Similar (Top) Hit BLAST E		5.3E-01	5.3E-01	5.3E-01	5.3E-01 /	5.3E-01 /	5.3E-01	5.3E-01	5.3E-01	5.3E-01		5.3E-01	5.3E-01	5.3E-01		5.3E-01	5.3E-01	5.3E-01					L		5.2			
	Expression Signal		6.83	2.74	1.58	1.96	1.96	0.84	0.84	1.82	1.82		8.	0.81	0.81		0.62	7.3	1.72	4.03	18.35	8 20							2.27
	ORF SEQ ID NO:		28231	28649		30860	30861	31158	31157	31266	1	1		34692	34603	l_		37480	37709		26229		26585			28521			3 28843
	Exon SEQ ID NO:		15491	15999	16928	18172	18172	1_	١	L	1		21494	21545	1	L	22759	24149	24379		L	Ì	13896	1	ı	1_	L	L	16193
	Probe SEQ ID NO:		2786	3237	4187	5371	5371	5466	5466	2	3	8000	8802	8854		\$0.88 80.88	10111	11550	11789	14877	767		1141	0100	2442	3447	3231	3400	3437

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Single Exon Probes Expressed in Brain	RF SEQ Expression (Top) Hit Acession Database ID NO: Signal BLAST E No. Source Source	0.78 5.2F.01 AETONOSE 4 No. Trucites dehydrogenase precursor (p1mdh) mRNA. nucites gene encoding	5.2E-01 6759047NT	5.2E-01 7108444 INT	-01 A4284281 4 EET ULIMAN	0.75 5.2E-01 X02218.1 NT	0.76 5.2E-01 X02218 1 NT	0.48 5.2E-01 AA194518 1 FET LINAN	5.2E-01 AF143952.2 NT		28013 1.84 5.1E-01 MRR800 1 NT LITTLE AND THE CONTROL OF THE CONTR	5.1E-011A.1933944 1	4.49 5.1E-01 A 1223044 4 NIT	5.1E-01 X87885 1 NIT	5.1E-01 BE883095 4 ECT LINVAN	5.1E-01 AI858405 1 EST LILLAN	2.81 5.1E-01 P96380 SWISSPECT	6.1E-01 U72653 1	5.1E-01 BF541068 1 FST UTIMAN	0.93 5.1E-01/AV712326 1 EST LIMAN	1.69 5.1E-01 R80873.1 EST HIMAN	0.83 5.1E-01 AW808881.1 FST HIMAN	0.63 5.1E-01 AW808881.1 EST HI IMAN	4.33 5.1E-01 J05412.1 NT	3.14 5.1E-01 W22302.1 FET LINAAN	0.89 5.1E-01 M94579.1 NT	6.1E-01 BF030207.1 EST HUMAN	5 1F-01 RE430089 4	1.24 5.0E-01 4886650 NT	1.24 6.0E-01
-		0		-										1.0	1.2												4.26	3,55		
\downarrow	ORF SEQ ID NO:			6						- -																			27690	27591
	SEQ ID NO:	16376	li	il	18364		1	1 1	22574	25010			13412	14394	14752		16904				- 1	ı	- 1		22240	22708	25137	24745	14861	14861
	Probe SEQ ID NO:	3823	456	495	5567	9830	9630	9832	9928	12744	603	633	633	1648	2017	4057	4164	5103	6128	6183	818	9470	8470	9583	9587	10060	12086	12326	2130	2130

Page 51 of 536 Table 4 Single Exon Probes Expressed in Brain

				Moet Similar		- I	
Probe SEQ ID NO:	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
							Buchnera aphidicola genomic fragment containing (chaperone Hsp60) groEL, DNA biosynithesis initiating buchnera aphidicola genomic fragment containing (gldA) genes, brotein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes,
	14870	27601	3.19	5.0E-01	01 AF008210.1	N	complete cds; and termination factor Rho (rho) gene>
	1						Buchnera aprildicola genomic inguirent Comming (****). Buchnera aprildedon protein (gldA) genes, protein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes.
2,5	14870	27602	3.19	5.0E-	01 AF008210.1	NT	complete cds, and termination factor Rho (rho) gene>
2014				5.0E	01 138483.1	LN.	Kattus norvegicus jagges protein marrial cas
3854	L	29241			-01 AB033010.1	뉟	Home sapiens minuted for the American Process Cone IMAGE:4271939 5
6547	1.		0.65	5.0E	5.0E-01 BF576199.1	EST HUMAN	A selvidonsis thallana DNA chromosome 4, contig fragment No. 49
7582	١_			5.0E	-01 AL161549.2	N.	Archidoseis theliane DNA chromosome 4, contig fragment No. 49
7562	1_	33335	0.75	5.0E	-01 AL161549.2		Areanopore amendia muscle beta-tropomyosin mRNA, complete cds
8428	l_				5.0E-01 M92304.1	N I	Achieved Roll MGC 79 Home sapiens cDNA clone IMAGE:4043485 3'
856	L		0.71	5.0E	5.0E-01 BF107848.1	ESI HUMAN	A 192200011 1111 AIGC 19 Homo saplens cDNA clone IMAGE:4136832 5
8358	1	33647	2.74	5.0E	-01 BF317212.1	ESI HOMAN	S SOCIAL DEBPANCHING ENZYME (GLYCOGEN DEBRANCHER) [INCLUDES: 4-ALPHA-
	1						GLUCANOTRANSFERASE (OLIGO-1, 4-1, 4-GLUCANTRANSFERASE); AMYLO-1, 6-GLUCOSIDASE GLUCANOTRANSFERASE
9525	5 22178	35362	1.36		5.0E-01 P35573	SWISSPROT	(DEXTRIN 6-ALPHA-U-GLUCOSIDAGE)) COCCEN DEBRANCHING ENZYME (GLYCOGEN DEBRANCHER) (INCLUDES: 4-ALPHA-
	i						GLICOGEN DELIGIONAL MANAGENERA SERVICE (DE CONTRANSFERASE); AMYLO-1,8-GLUCOSIDASE GLUCOSIDASE (DI COSIDASE))
9525	22178	35363	1.36		5.0E-01 P35573	SWISSPROT	(DEXI KIN 8-ALL HACL SECOND SERVICE ON A clone IMAGE:3849436 5'
10290	L	3	1.12		5.0E-01 BE869218.1	EST HOMAN	Min miscally MRC OX-2 antigen homolog gene, expns 2-5, and complete cds
12026	_	4	4	4 5.0E-0	5.0E-01 AF029215.1	z!	Home saplens chromosome 21 segment HS21C102
12715	ı	6	1.86		6.0E-01 AL163302.2	TOGGGGG	NUCLEAR ENVELOPE PROTEIN CUT11
12726					-01 013961	EST HIMAN	602076649F1 NIH_MGC_62 Homo sapiens cDNA done IMAGE:4243860 5
772	72 13544				4.9E-01 Brov 1402.1	LN	Xenopus laevis mRNA for c-Jun protein, 1978 BP
1656					4 1140980 4	LN LN	Cavia porcellus pulmonary surfactant protein A (SP-a) mRNA, complete cos
1899					4.9E-01 040008.1	SWISSPROT	FIBRILLIN 1 PRECURSOR
537	21 18124				4 4 700004 4	FN	Homo sapiens diacytalycerol kinase 3 (DAGK3) gene, exon 10
5946	46 18728				4.9E-01 AF020831.1	110	Homo sapiens diacy/glycerol kinase 3 (DAGK3) gene, exon 10
5946	46 18728				4.9E-01 AF020831.1	Į.	Oryza sativa subsp. Japonica mEF-G mRNA for mitochondrial elongation racid of continue was
7352	52 20033				4.9E-01 AB040031.1	TORGRENIA	PUTATIVE UNDECAPRENYL-PHOSPHATE ALPHA-N-ACETYLGLUCOSAWINTLI RANSFEDASE
7605		71 33378			4.9E-01 Q10000	SWISSPROT	PUTATIVE UNDECAPRENYL-PHOSPHATE ALPHA-N-ACETYLCLUCOSAMINTELINANS LINES
10/2	7605 20271				4.9E-01 010000	EST HIMAN	601874964F1 NIH MGC_54 Hamo sapiens cDNA clone IMAGE:4102503 5
88	8888 21579	79	1.45	١	4.8E-01 BF208/81.1		

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					S	ingle Exon Pr	Single Exon Probes Expressed in Brain
Probe		ORFSEO	Fxprescion	Most Similar	-	Hack	
NO.	NO:		Signal	BLAST E	I op Hit Acession No.		Top Hit Descriptor
9088	8 21775	34030	200				he90c02.x1 Soares NFL T GRC St Long State 1
9186			S. 60	4 BE-0	AW3390	EST_HUMAN	O95714 HERC2.; The septents conveyed and septents conveyed and septents of TR: O85714
10216	L	36076	28	4.86-01	4.9E-01 10946863 NT	3 NT	Mus musculus unc13 homolog (C. elegans) 1 (Unc13h1) mRNA
10419	L		12.0	4.80.4	4.9E-01 AF053980.1	LN.	Mus musculus adenyly cyclase 1 (Adcy1) cDNA, partial cys
11925	1		12.	4.95-0	1 X30000.1	L	H. sapiens DNA for BCL7A gene and BC 74/IGH locus fundamental
12709		1	2/:1	4.8E-0	E-01 AF176912.1	ż	Home saplens neurotrophin-//B-cell stimulating feater 3
	L	1	0.0	4.8E-0	4.9E-01 AA613562.1	EST_HUMAN	Inq22e11.s1 NCI_CGAP_Co10 Homo septems CDNA close 144 CE14110000
4288	17037		0.77	4.8E-01	4504850 NT	Į.	Homo sapiens potassium channel, subfamily K, member 5 (TASK-2) (KCNK5) mRNA, and translated
5420	18219	30830	10 78	1 0 10 7	10000		Saccharomyces cerevisiae) sporulation profein (SPO11) near coming of A
6229	l	32356	0.20	4.00-6	4.0E-01 JUZSBY.1	LN !	cds
6286	19352		3.76	A RES	-01 UB2002.1		Mus musculus slow skeletal muscle troponin T (Tnnt1) dene complete ods
7216	19901		188	7 20 7	/oggovy	EST HUMAN	nu85f09.s1 NCI_CGAP_Alv1 Homo sapiens cDNA clone IMAGE-1217613
7565		33339	0 78	4 8E-04	4 8E-04 AI 462200 C	z!	Home sapiens reproduction 8 (D8S2299E) mRNA
7661	20325	33434	4 05	A 8E 04	4.9E 04 A 464 400 0	N.	Homo sapiens chromosome 21 segment HS210009
7661	20325	33435	4.05	4 8F-04	4 8F-04 At 184402 2	Z	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 4
7805	00300				7501196.6		Arabidopsis thallana DNA chromosome 4, contig fragment No. 4
9144		33627	1.2	4.8E-01		EST_HUMAN	MER6 repetitive element :
10830	2000	-	0.92	4.8E-01	<u>.</u>	EST HUMAN	PM1+T0350-201209-004-604 LT0350-11
12247	25485	1	1.88	4.8E-01		N	S.cerevisiae ORFs from changes and a sapiens a DNA
12795	25218	1	3.04	4.86-01		ΝΤ	Trypanosoma cruzi transposon VIP II SIBE zazani
8422	19190	32186	1,00	4.8E-01			Chlamydomonas reinhardtil cop gene, exons 1-8
6841	19423	32438	200	4.75-01			601883880F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4062307 F1
7764	20460	33584	0 63	4.7E-01	01 71204374.1	T	q72a09.x1 Scares_testis_NHT Homo sapiens cDNA clone iMAGE-17555дд э
7764	20460	33585	0.63	77.0		٦	hbc811 Human pancreatic islet Homo sapiens cDNA clone hbc811 Fond
8974	21664	34816	0.52	4 7 1 0	04 8004 504	HOMAN	hbc811 Human pancreatic islet Homo sapiens cDNA clone hbc811 5end
10751	23438	-	6.11	4 7F.04	4 7E-04 A E400673 4		Kattus norveglous Spermine binding protein (Sbp), mRNA
11022	23694	36957	2.2	4 7F-01	T	N.	Influenza A virus isolate hk51697 hemagglutinin (HA) gene, partial cds
11252	23914	37206	1.61	4.7E-01		1444	Human collagen apha2(XI) (COL11A2) gene, exons 6 through 16, and partial cds
11349	24039	37342	1.7	4.7E-01.		7	002043888F1 NCI_CGAP_Bm67 Homo saplens oDNA clone IMAGE:4181303 6
12116	24609		1.52	4.7E-01 E	T	7	NOGAN I 0028-240400-011-E08 NT0029 Homo sapiens cDNA
12237	24689	\sqcup		4.7E-01 A	Т	Т	NOT 01 1335FT INIH _MGC_71 Homo sapiens cDNA clone IMAGE:3912488 6
					7		LI ICOO.XI SOBIES NFL T GBC S1 Homo sabiens china class (MAS) Francisco

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	t Top Hit Acession Database No. Source	1-2	04 AP-000007.1	041 B6/9502 NI	OT BESSOOT EST HIMAN	01 BF693300.1	04 BF313593.1 EST HIMAN	SIA CONTRACTOR	TORIGODA	CT CSUB45	Т	4.6E-01 Al247679.1 EST_HUMAN TR:015338 BUTYROPHILIN ;		SWISSPROT	LN	-01 BE817247.1 EST_HUMAN	4.6E-01 D26215.1 INT Markanobacterium thermoautotrophicum from bases 1165751 to 1176238 (section 100 of 148) of the	4.6E-01 AE000894.1 NT complete genome FIMPA (nempA) gene, mitochondrial gene encoding putative mitochondrial protein.	4.6E-01 U62332.1 NT complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds complete cds cds cds cds cds cds cds cds cds cds		4.6E-01 L07320.1 NT INUITIE COUNTING CAP Thy1 Homo sapiens cDNA clone IMAGE:943353 similar to contains Alu repetitive	4.6E-01 AA483577.1 EST HUMAN element.contains element.L1 repetitive element;	GENOME POLITING EN PROTEIN P3; 6 KD PROTEIN 1 (8K1); CYTOPLASMIC INCLUSION PROTEIN PROTEIN 1 (8K1); CYTOPLASMIC INCLUSION PROTEIN A (CI); 6 KD PROTEIN 2 (8K2); GENOME-LINKED PROTEIN (VPG); NUCLEAR INCLUSION PROTEIN A	SWISSPROT	4.6E-01 BF697399-1 EST_POWEN (GC-5) (4.8E-01 P55202 SWISSPROT CYCLASE)
-	Most Similar (Top) Hit BLAST E		4.7E-01 A	4.7E-01	4.6E-01 B	4.6E-01 BI	4.6E-01 BI	4.6E-01 BI	4.6E-01.0	4.6E-01 Q	4.6E-01 B	4.6E-01 A	4.6E-01 A	4.6E-01 P	4.6E-01	4.6E-01	4.6E-01	4.6E-01 A	4.6E-01	4.6E-01	4.6E-01	4.6E-01			4.6E-01	
	Expression Signal		1.83	1.38	1.57	1.57	-	-	3.11	3.11	48. 48.	2.17	2.17	9:	96.0	7.70	0.59	121	3.2	3.2	0.57	0.91		0.59	10.11	1.11
	ORF SEQ F				29116	29117	30795	30796	30875	30876	31148	31163	31164	1_			31526	31911	32620	32621	 _	33403	1	7	3 34049	0 35035
	SEQ ID		25055	25300	16479	16479	18136	18136	18185	18185	18258	18271	1	I	1	1	ı	18940	19586	1	1	L	1	20322	1	1 21870
	Probe SEQ ID		12813	12817	3726	3726	5333	5333	5385	5385	5459	5472		5480	75.00	5645	2888	6183	6989	999	7131	7870		7658	8219	9201

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ATRIAL NATRIURETIC PEPTIDE RECEPTOR B PRECURSOR (ANP-B) (ANPRB) (GC-B) (GUANYLATE Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds HUM105F03B Clontech human fetal brain polyA+ mRNA (#6635) Homo saplens cDNA clone GEN-105F03 w82e92x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2428618 3' similar to TR:Q92923 Q92923 SW/ISNF COMPLEX 170 KDA SUBUNIT.; POLY-BETA HYDROXYBUTYRATE POLYMERASE (POLY(3-HYDROXYBUTYRATE) POLYMERASE) (PHB POLYMERASE) (PHB SYNTHASE) (POLY(3-HYDROXYALKANOATE) POLYMERASE) (PHA POLYMERASE) (PHA SYNTHASE) (POLYHYDROXYALKANOIC ACID SYNTHASE) Homo sepiens hypothetical protein DKFZp547G183 (DKFZp547G183), mRNA Delinococcus radiodurans R1 section 68 of 229 of the complete chromosome 1 455402.s1 Soares, fetal, liver, spleen, 1NFLS, S1 Homo saplens cDNA clone IMAGE,484179 3* BASEMENT MEMBRANE.SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN PRECURSOR (HSPG) (PERLECAN) (PLC) wg73e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2370766 3° wg73e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2370766 3° Mus musculus DNA polymerase epsilon catalytic subunit (Pole) gene, exons 2 through 12 COLLAGEN ALPHA 5(N) CHAIN as96e09.x1 Berstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE;2363480 3° ho60g02.x1 Soares_NR_T_GBC_S1 Homo sapiens cDNA clone IMAGE;3041810 3° 601667225R1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE;3866023 3° Delnococcus radiodurans R1 section 68 of 229 of the complete chromosome 256g11,x1 NCI_CGAP_Ov35 Hamo sepiens cDNA clone IMAGE:2292844 31 Top Hit Descriptor Wg73e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saple PUTATIVE VITELLOGENIN RECEPTOR PRECURSOR (YL) L5+H70730-100500-075-g05 H70730 Homo saplens cDNA L5-H70730-100500-075-g05 H70730 Homo saplens cDNA QV2-PT0012-140100-031-c09 PT0012 Homo saplens cDNA D.melanogaster Shaw2 protein mRNA, complete cds Rat nucleolar proteins B23.1 and B23.2 Single Exon Probes Expressed in Brain COAT PROTEIN DNA PRIMASE EST_HUMAN SWISSPROT EST_HUMAN EST_HUMAN NT EST_HUMAN Top Hit Database SWISSPROT EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN HUMAN Source SWISSPROT EST_HUMAN SWISSPROT SWISSPROT SWISSPROT EST È Ę 11444786 NT Top Hit Acession 4.8E-01 BE186449.1 4.8E-01 AF019369.1 4.6E-01 AF019369.1 4.5E-01 Q28247 4.5E-01 AI708908.1 4.5E-01 AW873495.1 4.5E-01 BE983445.2 P98163 BE185449.1 AE001931.1 4.5E-01 AE001931.1 4.5E-01 AA677086.1 ģ 4.6E-01 A1915634.1 A1915634.1 AW608814.1 4.5E-01 AF128378.1 AI858849. 4.5E-01 AI848596. M37036.1 M32661.1 4.5E-01 Q52728 4.6E-01 4.6E-01 4.5E-01 4.6E-01 .5E-01 (Top) Hit BLAST E 4.6E-01 Most Simile 4.5E-01 4.5E-01 4.5E-01 <u>+ 8</u> 2.3 10.22 10.22 5.32 1.43 Expression 1.7 0.95 0.88 4.25 2.54 0.65 0.86 0.83 2.34 ORF SEQ ID NO: 35720 35721 36449 36850 36851 36450 27350 28722 28787 29431 33073 33273 33396 34292 34134 SEQ ID 22526 23602 23602 23217 14641 14641 15640 16072 16800 17617 ÿ 16131 17887 20287 21149 18260 19995 20900 21372 Probe SEQ ID 9876 10912 929 9876 10922 12163 11450 1904 3312 3372 4155 4890 4007 6510 7312 7609 7621 8206 8302 8457

Page 55 of 536 Table 4 SIndle Exon Probes Expressed in Brain

Single Exon Probes Expressed in Drain	Top Hit Descriptor	Escherichia coli K-12 MG1655 secton 108 or 400 or use conspice societical secton 108 or 400 or use conspice societical secton 108 or 400 or 108 or 10	Bombyx mori nuclear polymearosis winds, company general about clone HFBCY17	ES 102331 Fetal brain, Order 8 (cattle 36206) Homo saplens cDNA clone HFBCY17	CS102311 Start County County Homo sapiens oDNA clone IMAGE:2703985 3' similar to SW:INT6_MOUSE	XOI WILL IN TO STATION SITE PROTEIN INT-8. [1]; / QC4252 VILL INTEGRATION SITE PROTEIN INT-8. [1]; / QC4252 VILL PHOND septens dDNA done GLOCED12 5'	601449201F1 NIH MGC 65 Homo saplens cDNA clone IMAGE:3852961 5	602035275F1 NOL CGAP Brn64 Homo sapiens dDNA clone IMAGE:4183290 o	Homo saciens testis-specific kinase 2 (TESK2), mRNA	VASCULAR ENDOTHELIAL GROWTH FACTOR B PRECURSOR (VEGF-B) (VEGF RELATED	FACTOR)	Kartus norvegicus syrican is in variante in the syrican syrican is supplied to the syrican syr	Raftus novegicus Syrickin-5 million, 2011-100.	SA 22720E1 NIH MGC 44 Homo sapiens cDNA clone IMAGE:3609393 5	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	mirch frats. Sprague Dawley, sulfur-dioxde-freated tracheal epithelium, mRNA Fartal, 350 inj	AV720408 GLC Homo sapiens cDNA clone GLCCSC12 5	are 211 X NOT CGAP Brn 25 Homo sapiens cDNA clone IMAGE: 1861125 3' similar to I R. 4.29 I to 4.	UNKNOWN PROTEIN ; orichat.xt NGI OGAP_Bm25 Homo sapiens cDNA clone IMAGE:1861125 3' similar to TR:O29168 Q29168	UNIXNOWN PROTEIN:	ALATOXIN B1-ALDEHYDE REDUCTASE;	18780911.31 Strategic Complete LYN (HUMAN); TYROSINE-PROTEIN KINASE LYN (HUMAN); TYROSINE-PROTEIN KINASE LYN (HUMAN);	Heliconatcie pyon 2000 2000 Trenonema palijdum section 4 of 87 of the complete genome	S hiberosum mRNA for induced stolon tip protein (partial)	7/69a03.s1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:509639 3	HIV-1 isolate 08107/6 from USA, envelope glycoprotein (env) gene, partial cds	
e Exon Probe	Top Hit Database Source	LN LN		7	EST HUMAN		Т	TOT LINAN	N Carlo		ISSPROT		Т	EST HUMAN	EST HUMAN	SWISSPACI	SWISSLAND	NAME OF THE PARTY	בפו עמואסג	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN 1	I N	NAME OF THE PARK	FN - INCINION	
Sing	Top Hit Acession No.	4.5E-01 AE000218.1	30816					١	01 BF33/531.1	114220981N	01 P49765	01 AF058790.1	-01 AF058790.1	01 BF056726.1	-01 BE378707.1	-01 P04929	-01 P04929	01 S65019.1	-01 AV720408.1	4.4E-01 Al198413.1	4.4E-01 AI198413.1	-01 AW080795.1	4.4E-01 AA776132.1	4.4E-01 AE000571.1	4.4E-01 AE001188.1	4.4E-01 Z116/9.1	4.4E-01 AA056427.1	4.4E-01 AFT 16340.1
	Most Similar (Top) Hit BLAST E Value	4.5E-01	4.5E-01	4.5E-01 N	4.5E-01 M86006.1	4.5E-01 A	4.5E-01 A	4.5E-01	4.5E-01	4.5E-01	4.4E-01			4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4.4E-01	}	4.4E-01	4.4						
	Expression Signal	986	1.02	24.62	24.62	2.15	1.52	3.52	1.58	3.37	3.39	1.29	1,29	2.92	1.88	1.2	1.2	1.59	2	1.46	1.46	1.78	1.42	1.04	9.0	9.71		0.7
	ORF SEQ ID NO:	04770	37.5	36254	36255	1					77847			28723	L	30797	30798		31328	31591	31592	31894	1_	33056	6	9		34896
	SEQ ID	100	27,000	308	23038	l	L	<u> </u>	1_	24918	4 11400	Т	1	1	1_	ı	1	1	١.	18651	18651	L	l _		L.	20436		19 21738
	Probs SEQ ID NO:		688	10307	10392	10772	11217	11895	12540	12611	8	2000	3310	3343	4209	5334	5334	5602	5619	5864	5864	81.48	9 80	7297	7723	7740	8661	9049

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Single Exon Probes Expressed in Brain	It se Top Hit Descriptor		T	Т	7	Т.	Т	Deta -HKA=H,K-ATPase beta-subunit [rats, Genomic, 8983 nt, segment 2 of 2]	peta -HKA=H,K-ATPase beta-subunit [rats, Genomic, 8983 nt, segment 2 of 2]	wus musculus sodium channel, type X, alpha polypeptide (Scn10a), mRNA	T.	T	7	Callidrix Jaconus MW/LW opsin gene, upstream flanking region	7	T	Т	Turman Somatostatin I gene and flanks	Californ's Jacchus MW/LW opsin gene, upstream flanking region	California MW/LW opsin gene, upstream flanking region	Т	Т.	Т	Т	Columbia contraria large Rec. 2007 360 gene, partial cds	Family applies the state of the	T	Т	Т	Т	Equiple amy fourte may 2 and 1	HH744010 V NOT COATS	hh74e10.y1 NCI_CGAP_GL/I Homo septems cDNA_clone IMAGE::2968534 gr	TOTAL SERVICE SERVICE SOLVEN STONE IMAGE:2988654 5
Igle Exon F	Top Hit Database Source	į	ES HUMAN	SWISSPROI	EST HOMAN	SWISSPRO	OWING PRO	2 12	ž ž	E N	CMISSON	CAT LI MAN	NE LONAN		EST LIMBORY	EST CINAN	TIVE STATE	Ŀ			SIM/SCODE T	SWISSPROT	EST HUMAN	Į.	ĻΝ	Į.	SWISSPROT	ST HIMAN	L	E		EST HIMAN	EST_HUMAN	
ູ້ເຮັ	Top Hit Acesslon No.	4F-01 AWA12879 4	4 4F-04 OR2828	E-04 Albebeen 4	E-01 D28022	E-01 D25500	E-01 S78404 4	E-01 S78404 4	GRZZRZ4 NIT	TN 527749	1	4.4E-01/AW3R3338 1	ſ	T		T		ļ	ŀ	T	T					39.1		-01 BF348001.1				89		
	Most Similar (Top) Hit BLAST E Value	445-01	4 4 1 9	4 4F-04	4 4F-01		4 4F-01	4 4F-01	4.4E-01	4.4E-01	4.4E-01 P54725	4.4E-01	4.3E-01	4.3E-01	4.3E-01/	4.3E-01	4.3E-01 J00306 1	4.3E-01/	4.3E-01/	4.3E-01	4.3E-01	4.3E-01 F	4.3E-01 B	4.3E-01 A	4.3E-01 AJ001678.1	4.3E-01 A	4.3E-01 033367	4.3E-01 B	4.3E-01 U	4.3E-01/U	4.3E-01 Y	4.3E-01 A	4.3E-01 AV	
	Expression Signal	0.57	1.24	2.19	1,88	4.31	1.33	1.33	3.44	3.35	1.91	1.43	2.17	2.17	1.64	0.75	1.29	1.18	1.18	1.19	9.0	0.8	1.69	2.02	4.78	9.0	0.91	1.88	0.61	2.72	96.0	2.18	2.18	
	ORF SEQ ID NO:	34934	35025			35857	36132	36133	31095	31000			25835	25836		28466	29501	25835	25836		30742	30743	31515	31534	32384	32646			33260	1	35053	35468	35469	
	SEQ ID NO:	21771	21860	22612	22513		22821			_	24971	26152	13187	13187	15642	15822	16873	13187	13187	17629	18085	18085	18689	18000	188/1	9095	19511	20011	20168	21019	21885	22279	22279	
	Probe SEQ ID NO:	9082	9190	9862	9863	8997	10273	10273	12148	12679	12683	12766	402	402	2875	3056	4131	4374	4374	4902	6280	280	23/2	200	888	8	29/9	875	9	9770	8 8	920	9228	

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	Top Hit Acession Datebase No. Source	AW170669.1 EST_HUMAN	H65292.1 EST_HUMAN	AF075629.1 NT	EST_HUMAN	1 EST_HUMAN	EST_HUMAN	2.1 NT	SWISSPRO	NAMOL	Z	A1280338.1 EST_HUMAN	N81203.1 EST_HUMAN	1 Q04886 SWISSPROT	Injection (August 1): ST HUMAN III HISTOCOMPATIBILITY ANTIGEN, DR-1 BETA CHAIN (HUMAN);	EST HUMAN	A FST HIMAN	MANI ILL FOR	1 AW 854102.1	ALTOSZ47.2	AUTSS472.1	01 A0138472.1	01 S82504.7	1 AL101047.2	01 AW957448.1 ESI HUMAN	01 AW 957448.1 EST FOWER	01 4758039 NT	01 AA705007.1 EST HUMAN	01 AF181854.1 NT	01 AW863666.1 ESI HUMAN	01 AB023489.1	O1 BESO0483.2	
	Top Hit Database Source		T_HUMAN		HUMAN	Т	HOMAN	1000	STROI	NAMOI I		_	1	1		Т	Т	Т	7	_	_	Т	Į.	TOT LINAANI	TOT HOWAN	ES L'OIMEN	- 1				NOT TIMEN	FST HIMAN	
5	Top Hit Acesslo No.	AW170559.1	H65292.1	AF075629.1	AW993658.1	AW993658.1	AI874332.1	AJ003022.1	Q39102	AA761653.1	AE003947.1	AI280338.1	N81203.1	Q04886	A A 5340 Q3 1	D43487 4	DE040056 4	Drz42030.1	AW854102.1	AL163247.2	AU158472.1	AU1584/2.1	582504.1	AL10104/.2	AW957448.1	AW957448.1		AA705007.1	AF181854.1	1 AW863666.1	AB023489.1	1 BES00483.2	1 Viscostol:
	Most Similar (Top) Hit BLAST E Value	4.3E-01		4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.3E-01	4.2E-01	4.2E-01	4.2E-01	4.2E-01	4.2E-0	4.2E-01			1			4.25	4.25-		4.7	425	4.2E-	42E	4.2E	4.2E	4.2E	4. K	4.2E	4.2E	
	Expression Signal	48.0	0.5	2.45	1.29	1.29	1.84	2.18	1.64	1.23	4.4	1.41	0.85	76.0									2.15			2.21							3.5
	ORF SEQ ID NO:	888	36272	32646	37113		37662		26761			29021	L	28352									32694			33716	33932	L	35944				0 26488
	SEQ ID		22055	19606	1_	L	L	1	_	L	1	١.	1_	1_	١		4			18889			25101			1 20586	20800	1	L	_			2 13830
	Probe SEQ ID NO:		10128	10840	11166	11188	11745	12770	1337	1941	3596	3628	3698	3964		4649	4731	5626	5693	6112	6852	6852	6911	6993	7891	7891	810	9870	10081	10393	10972	11370	1072

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Campyobacter jejuni NCTC11168 complete genome; segment 3/6
AV649579 GLC Homo sepiens cDNA clone GLCBVD123*
PROBABLE SERINE PROTEASE DO-LIKE PRECURSOR (59 KDA IMMUNOGENIC PROTEIN) (SK69)
PROBABLE SERINE PROTEASE DO-LIKE PRECURSOR (59 KDA IMMUNOGENIC PROTEIN) (SK69) Mus musculus signaling intermediate in Toll pathway-evolutionarily conserved (Sitpec-pending), mRNA Voalavo gymnocaudus Vgym560 cytochrome b (cytb) gene, complete cds; mitochondrial gene for EST373364 MAGE resequences, MAGG Homo saplens cDNA Rhodococcus sp. AD45 isoG, IsoH, IsoJ, IsoA, IsoB, IsoC, IsoD, IsoE and IsoF gel om33d02.s1 Sogres_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1542819 37 ol94b08.st Scares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1505943 3' f002168590F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4297319 6' Methanococcus jannaschil section 77 of 160 of the complete genome 602133261F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4288238 6' VOLTAGE-GATED POTASSIUM CHANNEL PROTEIN KV1.1 (HUKI) (HBK1) Mus musculus platelet derived growth factor receptor, beta polypeptide (Pdgfrb), Delnococcus radiodurans R1 section 68 of 229 of the complete chromosome 1 Delnococcus radiodurans R1 section 68 of 229 of the complete chromosome 1 Mus musculus ubiquitin-protein ligase 63 componen n-recognin (Ubr1), mRNA Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36 EST373384 MAGE resequences, MAGG Homo saplens cDNA Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36 Top Hit Descriptor Homo sapiens anaphase-promoting complex subunit 7 (APC7), Laqueus rubellus milochondrion, complete genome Drosophila melanogaster Dalmatian (dmt) mRNA, complete cds Homo sapiens DNA for amyloid precursor protein, complete cds AV705243 ADB Homo sapiens cDNA clone ADBAHF08 6' CM2-HT0137-200999-010-e08 HT0137 Homo saplens cDN/ AV705243 ADB Homo saplens cDNA clone ADBAHF08 5 AV747880 NPC Homo sapiens cDNA clone NPCBDF10 5' Zea mays ZMPMS2 gene for 19 kDa zein prote Ascobolus immersus masc2 gene Ascobolus Immersus masc2 gene Single Exon Probes Expressed in Brain mitochondrial product EST HUMAN **EST HUMAN** EST_HUMAN Top Hit Database EST HUMAN HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN Source SWISSPROT EST_HUMAN NT SWISSPROT 6755521 NT F 6678490 NT Top Hit Acession 8404656 6679258 4.1E-01 AW961292.1 AL161536.2 4.1E-01 AL161536.2 4.1E-01 AL161536.2 4.1E-01 AV705243.1 4.1E-01 AV705243.1 4.1E-01 AW961292. 4.1E-01 AA909257.1 4.1E-01 AJ249207.1 4.1E-01 AL139076.2 4.1E-01 AV747880. BF681393.1 AF160597.1 4.1E-01 AV649579.1 BF349382.1 1.0E-01 AE001931.1 1.0E-01 AF203478. D87675.1 4.0E-01 Z96933.1 4.1E-01 X58700. 4.0E-01 Z96933.1 4.1E-01 Q09470 (Top) Hit BLAST E Vatue 4.1E-01 4.1E-01 dost Similar 1.1E-01 4.1E-01 4.16-01 4.1E-01 4.1E-01 4.1E-01 4.0E-01 4.0E-01 0.82 2.17 2.17 0.68 0.68 1.46 2.48 1.38 1.39 Expression 0.67 0.91 9.6 80.48 0.61 0.95 4.05 Signal 28161 29142 33092 ORF SEQ 29607 28706 31632 34827 36173 ÖNO 36673 36268 36404 27457 27619 27620 25595 13839 SEQ ID 15708 13839 16506 17353 16057 21678 16982 17011 18684 14215 15583 14886 2001 SEQ ID 3764 1081 2716 2941 2941 3754 4241 4868 4618 4271 7332 7935 8988 10310 10404 10743 10478 12476 1016 1468 1999 2156 2808

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			bp2x), and undecaprenyl-	se (mrar) genes, complete						COPROTEINS E3,					ıt																milar to TR:094821		æ
Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21C080	Homo sapiens chromosome 21 segment HS21C080	Streptococcus pneumoniae YilC (yilC), YilD (yilD), penicillin-binding protein 2x (pbp2x), and undecaprenyl-	pnospitate-UDF-imunyAc-pentapeptide phospho-munyAc-pentapeptide transferase (mrar) genes, compilete	10 GGG-/	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRDJ2), exon 1	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRDJ2), exon 1	NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN 5, CHLOROPLAST	EST382691 MAGE resequences, MAGK Homo sapiens cDNA	STRUCTURAL POLYPROTEIN (P130) [CONTAINS: COAT PROTEIN C; SPIKE GLYCOPROTEINS E3,	REAMORT, OND PETTIONS	MICRO LUBULE-ASSUCIATED PROTEIN 4	MR4-TN0110-180900-202-g02 TN0110 Homo sapiens cDNA	Homo sapiens OCTN2 gene, complete cds	EST26066 Cerebellum II Homo sapiens cDNA 5' end similar to EST containing Atu repeat	601558283F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3828092 5'	Synechocystis sp. PCC 9413 transposase gene, complete ods	Homo sapiens chromosome 21 segment HS21C100	HYPOTHETICAL 49.7 KD PROTEIN IN GIN2-STE3 INTERGENIC REGION	Gorilla gorilla carboxyl-ester lipase (CEL) gene, complete cds	Homo sapiens mRNA for KIAA1193 protein, partial cds	H.saplens B-myb gene	H.sapiens B-myb gene	Sinorhizobium meliloti egi, syrB2, cya3 genes and orf3	7161d01.x1 NCI_CGAP_Br16 Homo saplens cDNA clone IMAGE:3339169 3'	601563948F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3833639 5'	601862362F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:4082055 5'	Homo sapiens prepro dipeptidyl peptidase [(DPP-I) gene, complete cds	CM3-CT0105-170899-004-b08 CT0105 Homo sapiens cDNA	602019944F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4155322 5'	xn86d04.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2701351 3' similar to TR:094821	094821 KIAA0713 PROTEIN;	wp76a02x1 NG_CGAP_Bn25 Homo sapiens cDNA clone INAGE:2467658 3' stmilar to SW:RFX5_HUMAN P48382 BINDING REGULATORY FACTOR.;
Top Hit Database Source	NT	N P			Z.	Ŋ	N	SWISSPROT	EST_HUMAN	200000000000000000000000000000000000000	SWISSPROT	SWISSPROI	EST_HUMAN	NT .	EST_HUMAN	EST_HUMAN	NT	NT	SWISSPROT	NT	NT	NT	NT	된	EST_HUMAN	EST_HUMAN	EST_HUMAN	FN.	EST_HUMAN	EST_HUMAN		EST HUMAN	EST_HUMAN
Top Hit Acession No.	1 AL163280,2	4.0E-01 AL163280.2		, 0000000	4.0E-01 AF068903.1	4.0E-01 AJ277511.1	AJ277511.1	4.0E-01 Q31849	AW970610.1	20000	P2/285	P2/546	4.0E-01 BF092634.1	4.0E-01 AB016825.1	4.0E-01 AA323289.1	4.0E-01 BF030262.1	4,0E-01 L76080.1	4.0E-01 AL163300.2	P36049	3.9E-01 AF206618.1	3.9E-01 AB033019.1	3.9E-01 X82032.1	3.9E-01 X82032.1	3.9E-01 AJ225896.1	BF592611.1	3.9E-01 BE728667.1	3.9E-01 BF208036.1	3.9E-01 U79415.1	3.9E-01 AW177011.1	3.9E-01 BF348634.1		3.9E-01 AW195888.1	3.9E-01 Al937337.1
Most Similar (Top) Hit BLAST E Value	4.0E-01	4.0E-01	•		4.0E-01	4.0E-01	4.0E-01	4.0E-01	4.0E-01	10,	4.0E-01 P2/285	4.0E-01 P2/546	4.0E-01	4.0E-01	4.0E-01								3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01		3.9E-01	3.9E-01
Expression Signal	-	1.1			1.98	3.38	3.38	76.7	1.23		0.94	0,66	0.44	1.04	1.17	2.03	2.83	2.26	2.2	1.85	3.34	4.27	4.27	4.73	1.05	1.74	3.91	0.92	0.81	0.58		1.26	1.46
ORF SEQ ID NO:	L	28384			╽	29191			31538			33504	33649							26780				28499	29435	30270			Ì			34988	35301
Exan SEQ ID NO:	15734	l		0,0	16436	16559	16559	17499	18609		19116	20391	20524	20605	21595	24159	24316	25222	24972	14104	15358	15416	15416	15858	16804	17660	18631	20549	21452	21461		21822	22122
Probe SEQ ID NO:	2968	2968			3683	3807	3807	4767	2820		£43	7728	7829	7910	8904	11560	11721	12162	12684	1366	2648	2709	2709	3093	4029	4832	5843	7854	8760	8769		9134	9445

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Table 4
Single Exon Probes Expressed in Brain

									7	_	-	-	_	-	_	-	_	_	_	_		_,	_	_	_	_				_				_
Chigae Expire Expressed in Brain	Top Hit Descriptor	Himan clahindin 27 come commands and 44	Pornhuse appropries a feet and 11, and L1 and Alu repeats	Nicolana tahanim mDNA (x. 14.14.	AVR65874 GKC Home socions and Arts of the Complete cds	AV702823 ADB Home explains DNA class a Report of	Homo sapiens protocolven 3 (PPCs) some consists of	HOMEORY PROTEIN HI KA	Homo septens throughering profess (F) 10509 (F) 105001	Homo saciens profelin kinasa PKhibota (hizarioa), mkina	XVella fastidiosa section 16 of 220 of the complete contract	Ceanorhabditis bridgsae acetylcholinestorace (one 1) man and 101.	Arabidonsis theliana mitating man like terminal	Mus musculus solute carrier family 1 member 8 (SI-4-e)	Human immunodeficiency while the 4 complete control of the control	Pleuronectes emerican is emforment/deco N (2001)	Arabidonsis thelians DNA chromosom ()	W28H2 A Sorres NEI T CBO C4 L.	w/38h12x1 Source NET T GBC S1 Louis sapiens cDNA clone IMAGE:2367866 31	PM0-HT0339-200400-040-044 LT0330 Ltd. September Conversional IMAGE:2357866 3'	Mus musculus general franscrintion factor II ((GM3)) DNA	Takifugu rubribes wnt2 (partial) frank1 offr and frank2 (kerthal) seess	TRANSCRIPTION FACTOR SOX-10	prion protein [mink, Genomic, 2446 nt]	QV3-BT0537-271299-049-e02 BT0537 Homo sapiens cDNA	ha5411.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:2047917.3' similar to	Validancia theliana DNA ale	Maniscrittis gene fortoiling by the configuration of the configuration o	Willias bare for Kallikrein-binding protein	modes live receptor namangaus protein (LKH-1) mRNA, complete cds	Fromo sepiens mRNA for KIAA1631 protein, partial cds	namo sapiens FUS-like antgen-1 (FOSL1), mRNA	Homo septems chromosome 21 segment HS21C079 VE43h08 r1 Sognes fatal liver salesh 1NEI S. Homosomia	Afti repetitive element;contains PTR5 repetitive element;
שום באחוו הוכ	Top Hit Database Source	NT	ĻΝ	Į.	EST HUMAN	EST HUMAN	L	SWISSPROT	TN	LZ LZ	ŢN	N.	Į.	LN	Į.	LN	Z	EST HIMAN	EST HUMAN	EST HUMAN	Ę	TN	SWISSPROT	Þ	EST_HUMAN	EST LINAAN						E 4		EST_HUMAN
5	Top Hit Acession No.	-01 M19879.1	11465620 NT	D86722	3.9E-01 AV695974.1	-01 AV702623.1	-01 AF304354.1	Q61670	11433335 NT	7019488 NT	-01 AE003870.1	3.8E-01 U41846.1		3002	-01 AJ251057.1		T	Γ			34095	01 AJ271361.2			01 BE072399.1	01 At374601 1		T		-	1001	14 AI 482070 0	T	
	Most Similar (Top) Hit BLAST E Value	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01	3.9E-01 Q61670	3.9E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	.3.8E-01	3.8E-01	3.85-01/	3.8E-01 /	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01 A	3.8E-01 X	3.8E-01.N	3 RE-01	3 BE-04	3 BE-04 A		3.8E-01 T95413.1
	Expression Signal	3.03	0.58	0.77	1.98	1.47	3.37	208	1.44	8.33	1.03	1.29	1.62	3.98	1.14	1.39	7.98	0.79	1.22	1.15	0.97	0.74	1.42	0.74	5.5	4.58	1.25	4.42	0.86	204	102	1 28		3.55
	ORF SEQ ID NO:	35635		35932		37674						27918	28027	28092		28456	28887			29127	29287	29416	31221		32298	32614	32527		34028	34289	34358	34551		
	Exon SEQ ID NO:	22429	22496	22714	23410	24344	25295	24581	24891	12971	14801	15178	15290	15601	15769	15809	16233	16283	16283	16492	16647	16788	18320	19021	19294	19579	19502	20093	20890	21147	21215	21408		22011
	Probe SEQ ID NO:	9778	9845	10066	10722	11753	11948	12066	12559	158	1883	2460	2576	2638	3003	3043	3477	3527	3541	3739	3897	40 43	6522	8247	6258	6662	6840	7416	8186	8455	8523	8716		9461

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	Top Hit Descriptor	AV755814 BM Homo sapiens cDNA clone BMFBCE07 5'	RC0-H10841-040800-032-b12 H10841 Homo sapiens cDNA	y/92h11.s1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:30289 3'	y/92h11.s1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:30289 3	Borrella burgdorferl (section 10 of 70) of the complete genome	Human p53 (TP53) gene, complete cds	QV3-ET0063-190700-271-a05 ET0063 Homo sapiens cDNA	Mus musculus apoptosis inhibitor bcl-x (bcl-x) gene, exon 3 and complete cds	Mus musculus vomeronasal receptor V1rA4 (V1ra4) gene, complete cus	Mus musculus developmental control protein missy, parter	Homo sapiens mRNA for KIAA1410 protein, parda cos	Danjo rerio bone morphogenetic protein 4 precursor (BMr-4) gene, complete cue	ok39c07x1 Soares NSF F8 9W OT PA P S1 Homo sapiens cunk cone invace: 1310 106 3	MR3-OT0007-080300-104-b02 O10007 Hamo sapiens cLNA	Neisserta meningitidis serogroup B strain MC58 section 50 of 205 of the complete genome	Homo sapiens Interferon-Induced protein p78 (MX1) gene, compliste cas	Homo sapiens chromosome 21 segment HS21C0/8	Chicken (White leghan) delta-1 and delta-2 crystallin genes, complete ads	Mus saxicola haptoglobin mRNA, complete cds	Homo sapiens tumor endothelial marker / precursor (TEM/), mrNA	ya50a07,73 Soares fetal liver spiech TINFLS Homo sapiens culture invited 20224 3	hd45d05.x1 Soares_NFL_T_GBC_S1 Homo sapiens cUNA clone invAvEr.z912491 3 strities to contents rust repetitive element contains L1.12 L1 repetitive element;	Homo saplens chromosome 12 open reading frame 4 (C12ORF4), mRNA	Homo saplens chromosome 12 open reading frame 4 (C12ORT4), mKNA	okasb11.s1 NCI_CGAP_Let2 Homo sapiens cDNA clone IMAGE:1516/01 3	Gallus gallus mRNA for beta-carotene 15,15'-dioxygenase (bCDO gene)	mouse ig germline alpha membrane exons region	qt46b07.x1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:1930897.3	Rabbit mRNA for fast skeletal muscle myosin heavy chain (MHC)	Homo sapiens partial LIMD1 gene for LIM domains containing protein I and NIXXVO31 gene	Homo sapiens partial LIMD1 gene for LIM domains containing protein I and Nacross 1 gains	Bowne mKNA for terminal decynucieotrayiransiralase (1 d f / EC 2777.377)	UMZSUAVOD numar adia polyni (in girana) nomo odpore
ייי האיין וואס באו שניין וואס באו מייין	Top Hit Database Source	EST_HUMAN A	EST HUMAN R	Г	EST_HUMAN V	NT	H	T_HUMAN				NT		EST_HUMAN o		\neg						EST HUMAN y	EST HUMAN "			T_HUMAN			r_HUMAN				Т	EST_HUMAN
Sillo	Top Hit Acession No.	1 AV755814.1	1 BE719219.1			-		01 BE829256.1					3.7E-01 AF056336.1		-		01 AF135187.1	01 AL163278.2	01 M10806.1	10353.	11525843 NT	01 T66802.1	01 AW511326.1	11436739 NT	11436739 NT	01 AA902912.1	01 AJ271386.1	01 K00691.1	01 Al336411.1	01 X05958.1	01 AJ297357.1	-01 AJ297357.1	01 X04122.1	-01 D79348.1
	Most Similar (Top) Hit BLAST E Value	3.8E-01/	3.8E-01	3.8E-01 R42550.1	3.8E-01 R42550.1	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.8E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.75-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01	3.7E-01
	Expression Signal	1.67	3.18	2.27	2.27	4.76	2.08	3.39	1.54	1.74	1.51	12.24	9.64	7.39	1.3	2.55	1.15	6.0	0.66	0.72	3.23	9.0	0.56	2.07	2.07	0.65	1.31	0.6	4.12	1.98	2.81	2.81	2.75	1.43
	ORF SEQ ID NO:			37610	١						30966	27944									32794			34059		1			35970					
	SEQ ID	23386	24121	24288	24288	24636	25316	1	1	1	25040	ı	ŀ	1		1	١.	L	l.	1	L	1	l		1_		L	1_	L	L	23633	L		24271
	Probe SEQ ID NO:	10805	14534	11603	1168	12140	12270	12384	12723	12771	12788	2486	3453	4204	4786	4357	5678	7.860	6417	6436	7043	7685	77.10	8227	8227	8283	8 6	1000	10110	10764	10957	10957	11443	11676

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Single Exon Procession Detabase Source EST HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	PORMA I E HYDROGENLYASE SUBUNIT 5 PRECURSOR (FHL SUBUNIT 5) (HYDROGENASE-3 COMPONENT E) Homo sapiens PHEX gene 1/7/4808.r1 Soares fetal liver spleen 1NP.S. Homo sapiens ADMA Almo 11/4.05.7.7	y: 4500.1 Ocares four liver spreen TNFLS Homo septens cDNA clone IMAGE:275887 5' wt72Clox1 Soares_thymus_NHFTh Homo septens cDNA clone IMAGE:2813010 3' similar to TR:O16117 01617 FYN BINDING PROTEIN. [1]; SCO-SPONDIN
Source Source Source Source Source Source Source Source Source Source Source Source Source Source Source Source Source Source Source ST HUMAN INT INT INT INT INT INT INT INT INT IN	FORMA I E HYDROGENLY COMPONENT E) Homo sapiens PHEX gene vt74e06.r1 Soares fetal liver	wt72c10.xt Scares_thymus_NHFTh Ho O15117 FYN BINDING PROTEIN. [1]; SCO-SPONDIN
\$. \(\begin{aligned} \lambda \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\	SWISSPROT NT EST_HUMAN	HUMAN
		1.4
Most Similar (Top) Hit A BLAST E No Vellue 3.7E-01 J04982.1 3.7E-01 J04982.1 3.7E-01 J04982.1 3.6E-01 J05435.1 3.6E-01 J05655.1 3.6E-01 J056555.1 3.6E-01 J056	3.6E-01 P16431 3.6E-01 Y10196.1 3.6E-01 R94090.1	3.6E-01 A
Expression Signal Signal Signal 2.87 2.11 2.11 2.17 8.22 3.83 3.83 3.83 3.83 3.83 3.83 1.23 1.23 1.23 1.23 1.23 1.23 1.23 1.2	1.74	1.9
	31738	32943
	18778 19155 19739	19869
Probe SEQ ID NO: 11869 11284 1284 1284 1284 1284 1284 1284 128	7048 888 7048	7183

WO 01/57275 PCT/US01/00667

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		-		,—	-		_	_		_		_	,				_					_			_			
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 79	Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA	Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA	Homo sapiens chromosome 21 segment HS21C004	D. melanogaster singed gene, exons 3, 4, 5 & 6	D. melanogaster singed gene, exons 3, 4, 5 & 6	C.perfringens plc gene for phospholipase C upstream region containing bent DNA fragment	PROBABLE PEPTIDE ABC TRANSPORTER ATP-BINDING PROTEIN Y4TS	MR2-CT0222-211099-002-b10 CT0222 Homo sapiens cDNA	MR2-CT0222-211099-002-b10 CT0222 Homo sapiens cDNA	601676418F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3958997 5'	Arabidopsis thaliana mRNA for SigB, complete cds	Methandbacterium thermoautotrophicum from bases 702375 to 714311 (section 62 of 148) of the complete	Homo sapiens hHb5 gene for hair keratin exons 1 to 9	Synechocystis sp. PCC8803 complete genome. 3/27, 271600-402289	Escherichia coli K-12 MG1655 section 225 of 400 of the complete genome	Mus musculus Emr1 mRNA, complete cds	Homo saplens myelold/lymphold or mixed-lineage leukernia (trithorax (Drosophila) homolog); translocated to, 10 (AF10), mRNA	x80e11.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2679116 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	Mus musculus mannose receptor, C type 2 (Mrc2), mRNA	Homo sapiens GAP-like protein (LOC51306), mRNA	Homo sapiens GAP-like protein (LOC51308), mRNA	601811060R1 NIH_MGC_48 Home sapiens cDNA clone IMAGE:4053951 3'	601894653F2 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4124244 5'	Rattus norvegicus ADP-ribosylation factor-directed GTPase activating protein mRNA, complete cds	HOMEOBOX PROTEIN HOX-A4 (HOX-1.4) (MH-3)	zr08a09.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone IMAGE:650872.3'
gie Exon Pior	Top Hit Database Source	NT	LZ.	LΝ	LN LN	N _T	NT	LΝ	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	۲	5	Ę	Ę	Z	LN	N	EST HUMAN	N	N	NT	EST_HUMAN	EST_HUMAN	ΙΝ	SWISSPROT	EST_HUMAN
	Top Hit Acession No.	3.6E-01 AL161583.2	4504956 NT	4504956 NT	1 AL163204.2	1 X17550.1	X17550.1	3.6E-01 X62825.1	1 Q53194	3.6E-01 AW752901.1	AW752901.1	3.6E-01 BE902390.1	AB004293.1	3 6E-01 AE000856 1	3.6E-01 V19210 1	D90901.1	3.6E-01 AE000335.1	3.6E-01 U66888.1	11432598 NT	3.6E-01 AW190229.1	F 6678933 NT	7706136 NT	7706136 NT	3F129796.1	3.5E-01 BF310688.1	- -	506798	3.5E-01 AA223252.1
	Most Similar (Top) Hit BLAST E Value	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.65-01	3.6F-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01 U35776.1	3.5E-01 P06798	3.5E-01
	Expression Signal	11.45	2.74	2.74	1.17	1.04	1.04	0.57	14.67	0.51	0.51	3.31	4.12	3.4	1 83	4-	3.89	4	2.12	2.23	2.05	1.59	1.59	4.25	1.1	1.96	1.35	1.78
	ORF SEQ ID NO:	34005	34731	34732	34944	35139			35621	35735	35736	36791	36993	36419							25657	26131	26132	26194	27053	27069	27747	28066
	SEQ ID NO:	20871	21591	21591	21780	21986	21966	21944	22414	22543	22543	23544	23722	23188	25415	24522	24528	24624	24850	25363	13017	13482	13482	13535	14362	14382	15006	15600
	Probe SEQ ID NO:	8177	8900	8900	9091	9289	8239	9369	9763	9893	9893	10864	11052	11421	11903	11978	11987	12135	12493	12746	204	708	208	762	1615	1636	2281	2612

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Table 4

	Top Hit Descriptar		Sione IMAGE:1172357 3	e cds	S CLINA		c uguour			\$ NOTE OF THE PROPERTY OF THE	ZW79f03.r1 Scares_testis_NHT Homo sapiens cDNA clone IMAGE.782429 6' similar to TR:G1086935		GI I COSE-a PHOSPHATE 1 DELIVED CONTRACT OF STATE OF STAT	CHLOROPLAS PRECURSOR (G6PD)	10ASEV (LIEBES)	LIGASE) (FIGES)	EIGAGE) (FIGAG)	SOBP Z), mRNA	CONA	A, complete cds	VOLTAGE-DEPENDENT N-TYPE CALCIUM CHANNEL ALPHA-18 SUBUNIT (CALCIUM CHANNEL, L	N CALCIUM CHANNEL III) (BIII)	< A	V. Co.	p according	id exon 1	id excit i	V290h12.r1 Soares multiple sclerosis 2NhHMXO Dame contract Child Little Child	Sapiens CDIVA Glone IMAGE: 2803/6 6	exoll 1	V Z	tion of the state	genome
Single Exon Probes Expressed in Brain		preodos es NCI COAB V see Hand	Danio rotio homochous and Alexander Con Con IMAGE:1172357 3	RCS-HT0218-181090 011 02 LT0218 11	Mus musculus Alx42B gene 6' finding	Rat laukovita common antinan (1. CA) and	EARLY E24 DNA-BINDING PROTEIN	EARLY E2A DNA-BINDING PROTEIN	Himan mRNA for KIA Annae gong complete and	PM4-SN0012-030400-001-411 SN0012 Home stations about	Zw79f03.r1 Soares_testis_NHT Homo	Bos falinis partide mothinging autiend	GILLOSE - PHOSPHATE 1 DELIVER RECEISE (MS/A) mKNA, complete cds	S. scrofa mRNA for CD31 protein (PE)	HISTIDYL-TRNA SYNTHETASE (HISTIDINE TONA LIDASE) ALISES	HISTIDYL-TRNA SYNTHETASE (HISTIDINE—TRNA LIGASE) (HISTS)	Homo septens fumor profeso n.S. Ninding and an office of the profesor of the p	RC4ET0024-260600-014-007 ET0024 Home South - ENIA	Rattus norvariens Ne.KClockersconder (Allens)	Homo sabiens tyrosine kinase non-recentor 1 (TNK1) mBNA	VOLTAGE-DEPENDENT N-TYPE CA	X denvis dene for albitrata includire 1907 Onthe 30 (BRAIN CALCIUM CHANNEL III) (BIII)	QV2-HT0577-090400-128-07 HT0577 Homo sanion ability	C.griseus modopsin gene for onsin profein	Gallus gallus SPARC dene for osteonectin promotor and	Gallus gallus SPARC dene for osteopectip, promotor and exon	Human breakboint cluster region (BCB) gene complete and	vz90h12.r1 Soares multiple sclerosis	Drosophila melanogaster dual har protein (BerH2) man	Human glucokinase (GCK) gene reneat notworphism	HA0542 Human fetal liver cDNA library Homo sepiens and	B.taurus atpA1 gene for F(0)F(1) ATP synthase alpha-surbunit	Thermotoga maritima section 86 of 136 of the complete genome
gle Exon Pro	Top Hit Database Source	EST HIMAN	LN	EST HIMAN	LN	LN	SWISSPROT	SWISSPROT	NT.	EST HUMAN	EST HIMAN		SWISSPROT	TN	SWISSPROT	SWISSPROT	TN	T HUMAN	Ľ	7	TOGGSSIWS	Į.	Т	Т				T HUMAN			EST HUMAN		
Sin	Top Hit Acession No.	-01 AA642138.1	3.5E-01 AF071253 1	E-01 BE146585.1	3.5E-01 Y18477.1	M18349.1	3.5E-01 Q96687	-01 Q96687	3.5E-01 D42045.1	3.5E-01 AW863916.1			3.5E-01 024357				11448042 NT			7610	-01 002284		=		-01 AJ243178.1		-01 U07000.1		01 M82885.1		-		1.1
	Most Similar (Top) Hit BLAST E Value	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.55-01	3.5E-01	3.5E-01	3.5E-01	3.55-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01 P47281	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01	3,5E-01	3.5E-01	3.5E-01	3.5E-01	3.5E-01 X64565.1	3.5E-01
	Expression Signal	0.85	1.67	0.94	1.02	4.58	0.76	0.76	1.13	96.0	0.6	0.96	1.08	4.24	0.55	0.55	2.19	0.71	0.63	1.17	1.62	5.64	0.96	2.76	2.39	2.39	1,34	1.64	1.71	1.51	1.36	1.47	2.32
	ORF SEQ ID NO:		29596				30882	30686	31152		32070	32124	32338		33207	33208		33790		34662	35481	35644	35713	36564	36875	36876	37419	37499		37601			
	Exon SEQ (D NO:	16537	16972							18921	19085	19129	19331	19438	20118	20118	20665	20668	21059	21517	22288	22437	22517	23327	23625	23625	24108	24184	24216	24279	24367	24578	24676
	Probe SEQ ID NO:	3785	4231	4443	4627	4880	5251	5251	5462	6143	6314	6329	9999	9269	7441	7447	7970	7973	8366	8825	9836	9286	9867	10635	10946	10946	1505	11585	11619	11684	11776	12063	12214

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| Probe
SEQ ID
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 | 9112 | 9112 | 9321 | 8848 | 9346

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Page 67 of 536 Table 4 Single Exon Probes Expressed In Brain

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Page 68 of 536 Table 4 Single Exon Probes Expressed II

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Single Extra Trobes Expressed in Brain	Top Hit Descriptor	SOVE ARROPET NILL MCC FE LI	60447278871 NIH MOC 98 U	601472788T1 NIH MCC 69 Home sablens cDNA clone IMAGE:3876763 3'	CIRCI MEDADAZATE PRATEIN (2007)	CINCOMIST ON ZOUTE PROTEIN (CS)	Flexibacter literals our Brane for DNA gyrase is subunit, partial cds	1y84h01.x1 NCI_CGAP_Kid11 Home septens cDNA cione INAGE:2285809 3' similar to contains Alu repetitive element contains 1 transferior contains Alu	by84h01.x1 NCI_CGAP_Kid11 Homo sepiens of speciments, testings of similar to contains Alu repetitive element contains alement 1 reporting plants alement 1 reporting plants.	1040RE Limon ford house 1 1 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2	602140372F1 NIH MCC 48 House 2211 PRIOR SEPTEMBER OF THE TEGT	AU126115 NT2RP4 Home content of NA alone IMAGE: 4301800 6	AU126115 NT2RP1 Home saplens CONA clone in Larr-1000130 6	MITOGEN-ACTIVATED PROTEIN KINASE KINASE KINASE 1 (MAPK/ERK KINASE KINASE 1) (MEK KINASE 1/ MEKK 1)	CM2-ETD041-180500-187-440-ET0044-11	CM2-ET0041-180500-187-410-E10041-1-0m0 sapiens cUNA	28 STACL SOSTER fotal line Albeit 1904 Land	RC4-TN0077-250800-011-004 TN0077 Home septem cDNA clone IMAGE::297649 3	Homo sapiens high-mobility aroup phosphomolein (HMA) C) 2000	D.mauritiana Adh gene	D.mauritiana Adh gene	602070802F1 NCI_CGAP_Bm64 Homo sabiens cDNA close IMAGE 327355 F	hv51g02.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3176978 31	GALECTINS (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PROTEIN) (36 KD	LECTIN) (CARBOHYDRA I E BINDING PROTEIN 36) (CBP 36) (LAMININ-BINDING PROTEIN) (LECTIN L-29) (CBP30)	ob71g02s1 NCI CGAP GCB1 Homo sonians cDNA class MACE (2000 25 C)	Rhizolum leguminosanim svan plasmid API 6 II and 2000	Homo sapiens aldehyde oxidase 1 (AOX1) mBNA	Pyrococcus horikoshii OT3 genomic DNA, 287001-644000 nt mostling (272)	Raftus nonegicus EH domain binding protein Epsin mRNA, complete cds
וקום באטו דיוט	Top.Hit Database Source	FST HIMAN	EST HIMAN	EST HIMAN	TOGGRAMS	N PO	Ż	EST HUMAN	EST HUMAN	EST HIMAN	EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	EST HIMAN	EST HUMAN	EST HUMAN	Т	Г	Į.	Г	EST_HUMAN	EST_HUMAN		SWISSPROT	П				LN LN
5	Top Hit Acession No.	BF213873.1	3.3E-01 BE619650 1	01 BE619650.1	01 P05691	AB034233.1	-01 AB034233.1	01 AI628131.1	3.3E-01 AI628131.1	1 N85148.1	11 BF683954.1	AU126115.1	3.3E-01 AU126115.1	362925	3E828481.1	3.3E-01 BE828461.1		-					BE219351.1		P47953	AA806621.1	X07990.1	6598319 NT	AP000002.1	AF018261.1
Most Similar	Most Similar (Top) Hit BLAST E Value	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01	3.3E-01/	3.3E-01	3.3E-01 Q62925	3.3E-01	3.35-01	3.3E-01 N69866,1	3.3E-01	3.3E-01	3.3E-01 X63953.1	3.3E-01 X63953.1		3.3E-01 B		3.3E-01 P	3.3E-01 A	3.3E-01 X	3.3E-01	3.3E-01 A	3.2E-01 A
	Expression Signal	0.74	0;F	1.9	1.18	0.71	0.71	4.82	4.82	1.68	18.62	0.48	0.48	0.81	0.81	0.81	2.62	2.77	2.27	3.13	3.13	1.7	11.61		3.23	3.06	1.87	1.71	3.34	2.33
	ORF SEQ ID NO:	31417		·	31688	H	32652	32560	32561	33458	34295	34497	34498	34852	35203	35204	35244	35174		36554	36555		37147		37313		25453	37266		
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	SEQ ID NO:	5700	5856	5856	5947	6695	9699	62.89	6289	7682	8460	8859	8659	9012	9278	9278	242	9462	9894	10622	7002	Logo	1198		11317	11719	11741	11977	12676	4

Page 69 of 536 Table 4 Single Exon Probes Expressed in Brain

Probe NO: NO: 1139 1139 1136 1136 1136 1136 1136 1136		N N	Expression Signal 1.43 27.53 1.25 1.25 1.25 1.25 1.25 1.25 1.25 1.25	Most Similar (Top) Hit BLAST E Value 3.2E-01 3	Top Hit Acess No. No. 1,161561.2 250202.1 0,48624.1 0,48624.1 ANU957194.1 AL111655.1 BF203817.1 D10872.1 D10872.4 AF016494.1 AF016494.1 AR002359.1 AR002359.1	T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN T HUMAN	Arabidopsis thalians DNA chromosome 4, contig fragment No. 61 Fusarium pose virus 1 RNA2 putetive RNA dependent RNA polymerase gene, complete ods Fusarium pose virus 1 RNA2 putetive RNA dependent RNA polymerase gene, complete ods Nulgatis arcE-1 gene LACTOSE PERMEASE (LACTOSE_PROTON SYA/PORT) (LACTOSE TRANSPORT PROTEIN) S.carevisiae chromosome I reading frame ORF YBR172c EST389264 MAGE resequences, MAGD Homo saplens cDNA Homo saplens promyelotoyle loukamin ainc finger protein (PLZF) gene, complete cds Human splens promyelotoyle loukamin ainc finger protein (PLZF) gene, complete cds Human promyelotoyle loukamin ainc finger protein (PLZF) gene, complete cds Human MAGD (STM) mRNA Rabbit bela-like globin gene for anytamine N-acetyltransferase Homo saplens symplekin (SYM) mRNA Rabbit bela-like globin gene for anytamine N-acetyltransferase Homo saplens symplekin (SYM) mRNA Rabbit bela-like globin gene for anytamine N-acetyltransferase Hypophetic CAC (STM) MCD (STM) mRNA Rabbit bela-like globin gene cluster encoding the epsilon, ganna, admine gene gartal (STM) mRNA Rabbit bela-like globin gene cluster encoding the epsilon, ganna, admined genes Gozosi 1972F1 NIH MGC, 67 Homo saplens cDNA Gozosi 1972F1 NIH MGC, 67 Homo saplens cDNA Glardia intestinalis pyruvateriflavododin oxidoreductase and flanking genes Gozosi 1972F1 NIH MGC, 67 Homo saplens cDNA Gozosi 1972F1 NIH M
7755				3.2E	3.2E-01 AJ277661.1	Z	Rat ISO-atrial natriuretic factor gene, complete cds
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8 8	8268 20962	34103	13.76		3.2E-01 AL161574.2	NT	Arabidopsis thailana DNA chromosome 4, contig fragment No. 70
2		8					

Page 70 of 536
Table 4
Single Exon Probes Expressed in

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	2010 FEET CO. 1111 - 11	COLOSSISSIT IN MGC_57 Hamo sapiens cDNA clone IMAGE:4076627 5	De lessacuri NIH_MGC_5/ Homo sapiens cDNA clone IMAGE:4075627 5/	Demococcus radiodurans R1 section 152 of 229 of the complete chromosome 1	Orycoolagus cuniculus lg H-chain pseudogene, V-region (VH6-a2) gene, partial cds	Oryciologus cuniculus 1g H-chaln pseudogene, V-region (VH6-a2) gene, partial cds	Homo saplens chromosome 21 segment HS21C004	Human monocyte antigen CD14 (CD14) mRNA, complete cds	Homo saplens 6-phosphofructic-2-kinase/fructore-2 a kinatas-1-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-4-	Company of the state of the sta	norm septens o-prosphoructo-2 kinasefructose-2,6-bisphosphatase (PF2K) gene, exons 12 and 13	Dorrella Durgdorfert plasmid cp32-2, erpC and erpD genes, complete cds, and unknown genes	INVSTUD: X1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3181569 31	Homo sapiens gene for AF-6, complete cds	ES104/02 Fetal brain, Stratagene (cat#896206) Home sapiens cDNA clone HFBD724	Drosophila melanogastar taminin A (Lam-A) mRNA, complete cds	601507820F1 NIH_MGC_71 Hamo sapiens cDNA clone IMAGE:3809532 5:	ELONGATION FACTOR TU (EF-TU)	Homo saplens deoxycytidylate deaminase gene, complete cds	601275480F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3816748 F/	ye90h06.r1 Sogres fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:126051 6' similar to	HORD Carlons (71A A 24.7)	Homo serions (14 Aut 2)	History of Section 114 gate product (KIAA0174), mRNA	incolos, i Sceles JYTL 1 GBC S1 Homo septems cDNA clone IMAGE:2875391 3'	mus muscaus gene for Set/1 hr kinase KKIAMRE, exon 6	Delicus carota mkNA for transcription factor E2F (E2F gene)	Ayrete testidiosa, section 130 of 229 of the complete genome	nomo sapiens nepatocyte nuclear factor-3 alpha (HNF3A) gene, exon 1	HYPOTHETICAL PROTEIN HI1236	S.cerewistae chromosome XV reading frame ORF YOL141w	Musculus mkny for polycystin	From septens tilamin 2 (FLN2) gene, exons 10 through 22	1944 IT 3 Seares fetal liver spleen 1NFLS Homo sabiens cDNA clone IMA CE 1000007 E
Igie Exon Pro	Top Hit Database Source	EST UP HAARI	TOT TOTAL	TION LINE	E L	F.	L	N	Z	۲	-		7	ES TOWAN	Т	TOMAN	٦	٦	ISSPROT	П	EST_HUMAN (ENT LIMAN	Number of the second		T HIMAN	N CANO				100000	DY ISSUED IN			T CHIMANI	
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3	Most Similar (Top) Hit BLAST E Value	3.2E-01	3.2€-01	3.2E-01	3.2E-01	3.2F-01	3 2E-04	2 25 04	9.45-01	3.2€-01 /	3.2E-01/A	3 2F-01	3 2F-01 B	3 25-04	3 25 04 1	-13	3.2E-01 L07288.1	3.ZE-01 B	3.25-010	3.2E-01 L39874.1	3.2E-01 B	3.1E-01 R18051.1	3.1E-01	3.1E-01	3.1E-01 A	3.1E-01 AF	3.1E-01 A.	3 1F-01 AF	3 1E-01 AF	3 1E.01 PA			3.1E-01 AF	3.1F-01/Rp	
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<u> </u>	Exon SEQ ID NO:	21091	21091			21263		1	_	21737	21737	22644	22747	22858	23261	25317	25374	24871	24055	25354		15386	15632	16532	15630	15933	16637	17636	18190	18311	18312	18322	18478	26087	
	Probe SEQ (D NO:	8398	8398	8471	8571	8571	8962	8976		9048	9048	9894	10099	10210	10566	12010	12392	12624	12655	12712		2677	2702	2702	2862	3170	3887	4908	6390	5613	6514	5524	9899	6191	

Page 71 of 536 Table 4 Single Exon Probes Expressed in Brain

Single EXON Probes Expressed in Drain Top Hit Deacriptor Source Source Source Source Source EST_HUMAN RC3-HN0001-310300-0011-b04 HN0001 Homo seplens cDNA EST_HUMAN RC3-HN0001-310300-0011-b04 HN0001 Homo seplens cDNA clone MAGE:3640420 of EST_HUMAN G01306121F7 NIH_MGC_38 Homo seplens cDNA clone MAGE:3640420 of EST_HUMAN G01306121F7 NIH_MGC_38 Homo seplens cDNA clone MAGE:36309 3' EST_HUMAN G0124745F NIH_MGC_38 Homo seplens cDNA clone MAGE:36309 3' EST_HUMAN G0124745F NIH_MGC_38 Homo seplens cDNA clone MAGE:428161 f 5' EST_HUMAN G0124745F NIH_MGC_38 Homo seplens cDNA clone MAGE:428161 f 5' EST_HUMAN MO154745F NIH_MGC_38 Homo seplens cDNA clone MAGE:428161 f 5' EST_HUMAN G0124745F NIH_MGC_38 Homo seplens cDNA clone MAGE:428161 f 5' EST_HUMAN MO165 \$1 Stategens felal spiene (#837205) Homo seplens cDNA clone MAGE:428161 f 5' EST_HUMAN MO1850-MAZE HAN G01C L081V GANA-A AND GAMAA-C CHAINS (HUMAN) to gab/M81036 maz HEMOGLOBIN GAMAA-A AND GAMAA-C CHAINS (HUMAN) to gab/M81036 maz HEMOGLOBIN GAMAA-A AND GAMAA-C CHAINS (HUMAN) to gab/M81036 maz HEMOGLOBIN GAMAA-A AND GAMAA-C CHAINS (HUMAN) to gab/M81036 maz HEMOGLOBIN GAMAA-A AND GAMAA-C CHAINS (HUMAN) Homo seplens RAMAA-C GAMAA-C GAMAA-C CHAINS (HUMAN) to gab/M81036 maz HEMOGLOBIN GAMAA-A AND GAMAA-C CHAINS (HUMAN) homo seplens RAMAA-C GAMAA-C GAMAA	Corynebacterium sp. ALY-1 alyPG gene for polyguluronate lyase, complete cds PMH-ST0262-261199-001-901 ST0262 Homo sapiens cDNA	Balaenoptera physalus gene encoding atrial natriuretic peptide 601594960F1 NIH MGC 9 Homo sepiens cDNA clane IMAGE:3948734 5'	Cantagalo orthopoxins hemagglutinin gene, complete cds
TOP Hit Source Source Source HUMAN R HUMAN Y HUMAN Y HUMAN CHUMAN	MAN	T - T	ESI TOMAN
88 121 12 12 12 12 12 12 12 12 12 12 12 12	-01 AF237778.1 -01 AB030481.1	_	3.0E-01 BE741629.1 3.0E-01 AF229247.1
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Strongdycentertus mineral conference | Congressionations purpured 3467 KDa laminin-binding protein mRNA, partiel cds
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 | Steplococolis pre impolae strain DBI E D | Thermotoga maritima section 67 of 136 of the complete access | Mus musculus C-type (calcium dependent, carbohydrate recognition domain) lectin, superfamily member | (VIOLESIA), IMININA
604330070E4 NILL 1100 TO 11
 | Street NIH MGC 53 Home sapiens cDNA clone IMAGE:3681594 5 | Surpriorityces surrondraciens isopenicillin N synthase (pcbC) gene, partial cds | ronno sapteris DKFZP300M/122 protein (DKFZP586M0122), mRNA
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 | RCZ-BN0074-240400-110-h12 BN0074 Homo saplens cDNA | Social 2027 11 TINH_MGC_81 Home septens cDNA clone INAACE:4288336 5' Actinobecillus estinomycetemcomittens TadA (tadA), TadB (tadB), TadC (tadC), TadD (tadD), TadE (tadE), TadE (tadE) and TadC (tadD), TadE (tadE) | Actinobacillus actinomycetemcomitens TadA (tadA), TadB (tadB), TedC (tadC), TadD (tadD), TadE (tadE) | radii (kaur /), and i add (kadd) genes, complete cds

 | Specially souzes big and for EP changes in the september of the second state of the second se | 02140133F1 NIH MGC 46 Home sealers CDNA - I WOOF GO | 102140133F1 NIH MGC 46 Homo saplens CDNA clone IMAGE:4201097 5
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 | hrysodidymus synuroideus mitochondrion complete genome | PM1-CT0326-171299-001-f12 CT0328 Homo saplens cDNA |
| Top Hit
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 | | 2.9E-01 A |
| Expression
Signal | 3.94 | 3.94 | 4.77 | 3.06 | 0.61 | 0.85 | 0.71 | 2.77
 | 1.37 | 1.3 | 2.97 | 1.32
 | 0.51 | 8.0 | 200 | 43.84

 | 1.25 | 0.49 | 0.49 | 0.84

 | 1.88 | 0.73 | 0.73
 | 1.95 | 1.95 | 2.62 | 5.51 | 1.43
 | 1.16 | 2.73 |
| ORF SEQ
ID NO: | 30924 | 30925 | | | 30567 | 32510 | |
 | 33159 | 33648 | | 34207
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 | 35714 | 35908 | 35909 | 36155

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| Probe
SEQ ID
NO: | 5417 | 5412 | 5453 | 6732 | 6762 | 6827 | 954 | 122
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8 | 7827 | 8271 | 8374
 | 8728 | 8770 | 9118 | 9826

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| | Exon ORF SEQ Expression (Top) Hit Top Hit Acession No. Signal BLASTE No. Source | Exon SEQ ID NO: Expression 10 NO: Recession 10 NO: Signal No: Place No: Top Hit Acession No: Top Hit Acession Source Top Hit Acession Source Top Hit Acession No: Source Source 18216 3.0924 3.06-01 BE683576.1 FST HIMAAN RC3-BT0333-16077614.1 RC3-BT0333-16077614.1 | Exon SEQ ID NO: 10 NO | Exon NO: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Acession No: Top Hit Acession Source Top Hit Acession Source | Exon ORF SEQ Expression 10 NO: Signal (Top) Hit Acession 10 NO: Signal Top Hit Acession 10 NO: Signal Top Hit Acession 10 NO: Signal Top Hit Acession 10 NO: Signal 10 NO: Source 10 No: | Exon
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BE693675.1 EST HUMAN 18252 31442 4.77 3.0E-01 DI01247.1 NT 18253 3.0E-01 DI03249.1 NT NT 19366 3.0E-01 DI03269.1 NT 19489 3.0E-01 AF229247.1 NT 19713 3.2E-01 AF071810.1 NT 20078 3.3159 1.37 3.0E-01 AF071810.1 NT 20522 3.3648 1.3 3.0E-01 AF071810.1 NT 20966 2.97 3.0E-01 AF071810.1 NT 2052 3.3648 1.3 3.0E-01 AF071810.1 NT 20966 2.97 3.0E-01 AF0 | Expn
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	Expression Signal		2.73	0.72	0.73	7.70	1.1	1.66	0.79	5.19	5.19	6.4	1.47	1.08	2.4	0.58	0.58	0.56				2.00				0.48	9.0		0.49
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Page 74 of 536 Table 4 Single Exon Probes Expressed i

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Sirigle Exon Probes Expressed in Brain	Top Hit Descriptor	NI LEPONO III	AUTSUSTO NIZKPZ Homo sepiens cDNA clane NTZRP20039013*	Arabidopsis thaliana sulfonylurea receptor-like protein mRNA, complete cds	Baboon lymphocyte homing/adhesion receptor mRNA, complete cds	Pyrococcus abyasi complete genome; segment 5/6	Pyrococcus abyssi complete genome; segment 5/8	I rypanosoma cruzi stage-specific surface glycoprotein gp82 (gp82) mRNA_partial cds	l orpedo californica mRNA encoding acetylcholine receptor gamma subunit	ny35h02.s1 NCI_CGAP_Pr12 Homo saplens cDNA clone IMAGE:12737770 clossing	repetitive element;	Campylobacter jejuni NCTC11168 complete genome; segment 5/6	Kattus norvegicus activin receptor-like kinase 7 (ALK7) mRNA. compete cds	Rattus norvegicus activin receptor-like kinase 7 (ALK7) mRNA, complete ces	Homo saplens TNF-a-induoible RNA binding protein (TIRP) gens, complete cds	Chlamydomonas reinhardtii mRNA for nitrite reductase structural forus	Chlamydomonas reinhardtii mRNA for nitrita reductase structural locus	Rattus norvegicus A-kinase anchoring protein AKAP150 mRNA, complete cals	Prune dwarf virus movement protein, complete das coat protein, complete cds	Guira guira oocyte maturation factor Mos (c-mos) gene, partial cds	601148733F1 NIH_MGC_19 Homo sepiens cDNA clone IMAGE:3163688 5	001146/33F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3163688 gr	numan mkny for senne/threaning protein kinase, complete cds	Treparus G Wrus Isolate 80 (SZNAE12) polyprotein precursor, gene, partial cds	CV 1-010304-12020U-063-505 C10364 Homo sapiens cDNA	131 CONTROL STATE OF	Fesherichia Ali V. 40 MS 200 S1 Homo sapiens aDNA clone IMAGE: 2912333 3'	Sometimes can R-12 MG1855 Section 384 of 400 of the complete genome	Solds will a can K-12 MG1855 section 384 of 400 of the complete genome	Arekardopsis indigene UNA chromosome 4, contig fragment No. 65	Arabidopsis thaliana mRNA for lipoyitransferase, complete cds	l oxopiasma gondii 90kDa heaf-shock protein (HSP90) mRNA, partial cds	B taring microsateline (ETH121)	o. dan us microsatelitie (E.1.H121) Pyrocociiis hydiosali 072	Coccession of Spenomic DNA, 777001-994000 nt. position (4/7)
gie Exon Pro	Top Hit Database Source	For university	NAMOU TO THE	Z	Z	Z			Z	Т	HOMAN	Ž									HOMAN	TIONAIN DE		HIMAN	Т	Т	NEWIOL -								
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	Expression Signal	0.84	1.02	0.65	0.78	0.78	1.93	1.75	1.75	1,50	3.55	1.62	1.82	4.05	1.86	1 86	1	101	3.89	1.62	1.62	1.34	1.11	2.04	2.35	1.41	2.04	2.04	1.89	26.0	1.29	2.04	2.04	1.13	
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Page 75 of 536 Table 4 Single Exon Probes Expressed in Brain

Single Exon Probes Explesses in 2000.	Most Similar Top Hit Acession Top Hit Square Signal PLASTE No. Source	L. Constant	1.59 2.8E-01/AE001180.1	1.95 2.8E-01 A1090868.1 EST_HUMAN	0.99 2.8E-01 AL021127.2 NT	2.31 2.8E-01 P13615 SWISSPROT	1.19 2.8E-01 AF075238.1 NT	2.67 2.8E-01 AF030154.1	1.23 2.8E-01 BF528188.1 ES1_HOWEN	1.91 2.8E-01 AI272669.1 EST HUMAN	24.65 2.8E-01 AA349997.1 EST_HUMAN	3.07 2.8E-01 AB016625.1 N1	EST HOWEN	1 EST_HUMAN	2.8E-01 M3668.1 NT	1.53 2.8E-01/AF003124.1 NT	1.63 2.8E-01 AF003124.1 NI	8.34 2.8E-01 BF511215.1 EST HOME	TN	0.58 2.8E-01 X69980.1 N1	1.28 2.8E-01 AI346128.1 EST_HUMAN	1.28 2.8E-01 AI346126.1 EST_HUMAN	1.92 2.8E-01 U51688.1 NT	EST_HUMAN	7.34 2.8E-01 BF347847.1 EST HUMAN	Ę.	0.88 2.8E-01 L13654.1 NI	
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SEQ IO NO:	SEO ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acess	ngle Exon Pro	Single Exon Probes Expressed in Brain Top Hit Database Source
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865	7 21349	34494	0.59		014764	N.	Arabidopsis thallans DNA chromosome 4, contig fragment No. 52
8925	,	34760	0.48		X03246 1	SWISSPRO!	MAJOR VAULT PROTEIN (MVP) (LUNG RESISTANCE-RELATED PROTEIN)
923.	١,	35084	10.41	1	ORSBOD	NI CONTROL	Staphylococcus aureus transposon Tn554
9237	ı	35085	10.41	2.7E-01	083809	SWISSPRO!	THREONYL-TRNA SYNTHETASE (THREONINE-TRNA LIGASE) (THRRS)
823	1		2	2.76-01	2.7E-01 P37928	SWISSPROT	I HKEONYL-TRNA SYNTHETASE (THREONINE-TRNA LIGASE) (THRRS)
9700		35546				DALESCA	Rattus norvegicus DNA for peroxisome accomplisation
998	22629	35838	0.01	2.7E-01	2.7E-01 D89660.1	N	and complete cds
10018	1	35882	7.00	2.75-01	AF091848.1	ΝΤ	Oryctolagus cuniculus calgranulin C mRNA, partial cde
10149	22797	36012	3.08	2.75-01	AF087434.1	LΝ	Mus musculus transcription factor NF-A To Isoform a (NF-A Ton) - ISNIA
10149	1	36013	0.00	2.75-01	1	NT	Homo sapiens xeroderma pigmentosum complementation are a 2000000
10714	L	26645	0.0	2./E-01		L'N	Homo sapiens xeroderma planentreum complementation group C (AFC) gene, Intron 9
10714	1	36642	1.62	2.7E-01	2.7E-01 AV705043.1	EST_HUMAN	AV705043 ADB Homo saplens cDNA clone a DBCODOR # 1
	ı	1	70.1	2./E-01	AV705043.1	EST_HUMAN	AV705043 ADB Homo saplens cDNA clone ADBCODOS
10724	_	36653	3.13	2.7E-01		ŀ	Homo saplens caveolin-1/-2 locus, Contig1, D75622, genes CAV2 (exons 1, 2e, and 2h) CAV4 (
11820			1.5	2 7E-01	T	T.	CALL STATE OF THE
12482			1.49	27F.01	T	SOFKO	PUINTIVE 60S RIBOSOMAL PROTEIN C4F8.05C
12665	24963		2.83	2.7E-01	Τ	E A	Arabidopsis thaliana mRNA for sulfate transporter, complete cds
12811	25054	30929	1.95	2.7E-01	1	7	nomo sepiens fragile 16D oxido reductase (FOR) gene, exon 6
457	16542	25883	2.03	2.8E-01	2.6E-01 P78411	EST TOWAY	AV 42419 CB Home saplens cDNA clone CBMAXF02 5'
468	13254		1.38	2.6E-01	1	┪	INCAUCIS-CLASS HOMEODOMAIN PROTEIN IRX.2
1372	14120	26795	1.65	2.6E-01	T	111111111111111111111111111111111111111	bos taurus mRNA for mb-1, complete cds
1417	14165	26848	1.14	2.6E-01	T	Т	ou is i usa 84-1 NIH _MGC_71 Homo saplens cDNA clone IMAGE:3912346 67
1889	14626	27335	4.33	2 RE-01 A	T		Glycine max pseudogene for Bd 30K
1889	14626	27336	7 33	10.0	1		Arabidopsis thallana DNA chromosome 4, contra framment No. 2
		+	30.7	2.05-01/	U1 AL161472.2	/	Arabidopsis thaliana DNA chromosome 4. contin fromment No. 2.
				<u></u>		<u></u>	bb04d10.x1 NIH_MGC_14 Homo sapiens cDNA clone IMAGE:2058464 2: -1
2086	14818	_	10.48	2.6E-01/A	01/AW733152 1		RIBOSOMAL PROTEIN L7A (HUMAN); gb:M14889_cds1 Mouse surfelf focus surfelf 3 profeth games
2148	14878	27613	3.7	2.6E-01 N	T	T	
2476	15194		1.62	2.6E-01 Y			Trumen preaibumin gene, complete cds
							O. Maritanus rock gene

Page 79 of 536 Table 4 Single Exon Probes Expressed in Brain

Single Exon Flubes Expressed in Orani	Top Hit Descriptor	801128018E1 NIH MGC 9 Homo sapiens cDNA clone IMAGE:2890043 5'	Becterinhage T2 DNA-(adenine-N6)methyltransferase (dam) gene, complete cds		Homo sapiens acetylcholinesterase collagen-like tail subunit (COLQ) gene, exons 1A, 2, 3, 4, and 5	EST371580 MAGE resequences, involvintum capture of the contraction of	QV1-B10630-040400-132-eux B10630 Harro Capacita	Enterocococus racement or an incomplete cost and the service of the complete cost and the service of the cost of the service of the cost of the service of the cost of the cos	Salius galus III v.Y. is sweletal mosin heav chain, complete cds	Callus galds III vo 19 2000 1000 1000 1000 1000 1000 1000 1	sacodo 7.1 Stranggoro 1000 (Lhos 11 chlorophyll afb-binding protein (Lhos 3-1) mRNA, complete cds	Argundusis demand of the management of the manag	Ophrestia radicosa maturaso-like protein (matk) gene, complete cds; chloroplast gene for chloroplast product	Mus musculus metalloproteass distribution (North Programme) (North	Money in Series produce for PAP, complete ods	Taranicum Code Code Homo saplens cDNA clone IMAGE:2075788 3' similar to contains element	MERS5 repetitive element: MERS5 repetitive element: IMP-calactose translocator, pim-2 protocnoogene homolog	Homo saptens protein translocated, which protein is a protein and transcription factor IGHM inim-2h, and shal-type potassium channel genes, complete cds; JM12 protein and transcription factor IGHM	enhancer 3 genes, partial cds; and unknown 9>	I nermotoga intariorità coccari	ts02e12.x1 NCI_CGAP_Pan1 Homo sapiens cDNA done IMAGE:2227438 3' simitar to SW:NDF1_KAI 1084289 NEUROGENIC DIFFERENTIATION FACTOR 1; contains element LTR1 repetitive element;	ts02e12.x1 NCI_CGAP_Pan1 Homo sapiens cDNA done INAGE:2227438 3' similar to SW:NDF1_RAT Q64289 NEUROGENIC DIFFERENTATION FACTOR 1 ;contains element LTR1 repetitive element; Q64289 NEUROGENIC DIFFERENTATION FACTOR 1 ;contains element 6/7	Nelsseria meningitidis serogroup A strain 12-49 Comprose Services	601581754F1 NIH_MGC_7 Homo saptens cUNA cione intrace: 2015	601581754F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3938130 3	w448c04.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone liviading the liviading to grant and saplens control in the livian in grant and saplens control in the livian in the livian in the livian in the living control in th	7
le Exon Pion	Top Hit Database Source	NAME	NOW	Z	N	EST_HUMAN	EST_HUMAN	LN TN	LN	LN	EST HUMAN	Ł	LN.	N	EST HUMAN	Z	EST_HUMAN		LN LN	LN.	EST_HUMAN	EST_HUMAN	N.	EST HUMAN	FST HUMAN		EST_HUMAN
Sing	Top Hit Acession No.		1 BE272440.1	11 M22342.1	2.6E-01 AF229118.1	2.6E-01 AW959510.1	11 BE080598.1	01 AF175293.1	01 AB021180.1	2.6E-01 AB021180.1	2.6E-01 AA457617.1	01 U01103.1	.01 AF142703.1	01 AF153350.1	-01 H04858.1	-01 AB035972.1	-01 AI862398.1		2.6E-01 AF207550.1	AE001811.1	-01 Al682557.1	2.6E-01 Al582557.1	2 RE-04 Al 162757.2	2.3C-01 RE702052 1	04 BE702052 1	DE LOCACE.	2.6E-01 AI914380.1
,	Most Similar (Top) Hit BLAST E.	ŧ	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01		2.6E-01	2.6E-01	2.6				20.2	P. P.	1
	Expression Signal		8.87	8.99	2.02	98.0	16.7	1.2	8.0	8.0	1.46	1.77	1.18	0.95	3.6	1.06	0.68		0.73	235	2.26					0.73	0.9
	ORF SEQ ID NO:			28971	29028	20020	29506	29691							30257				31400		31854		١			4 32345	32719
	SEQ ID		15263	16323	16297	1	1	1	١.	١	1_	1_	1	1	1_	١.	<u> </u>	_	18481	1.	4000 A		- 1	- 1		0 19334	8 19673
	Probe SEQ ID		2548	3568	180	300	4078	4324	4459	4459	4511	4601	100	4007	4914	5257	5484		6883	2980	200		6108	6328	6570	6570	8669

Page 80 of 536 Table 4

			T		П		T	Τ	T	T	Τ	T	T	Т	Т	Т	Т	_	Γ	Т	Т	Т	П		Т	7	7	7	_		_	
Single Exon Probes Expressed in Brain	Top Hit Descriptor		CM0-H10245-031199-085-f04 HT0245 Homo sapiens cDNA	Campylobacter jejuni NCTC11168 complete genome; segment 4/6	2992801.1.1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone IMAGE:827672 6: y137803.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:1200A 32 cleating to	195:X12517 U1 SMALL NUCLEAR RIBONUCLEOPROTEIN C (HUMAN):	HYPOTHETICAL TRP-ASP REPEATS CONTAINING PROTEIN C29E8.01 IN CHROMOSOME	Jeografi To Soares fetal liver spleen 1NFLS Homo sapiens oDNA clone IMAGE:124212.5'	Mr.C-H 10166-181199-003-d12 HT0168 Homo saplens cDNA	Umelanogaster mRNA for alpha 1,2 mannosidase (Berlin)	D. meianogaster mKNA for alpha 1,2 mannosidase (Berlin)	UVBOTUTE INCL. CC3AP Brn64 Homo sapiens cDNA clone IMAGE:4150396 5'	POE ETABLE 10AL 75.2 KD PROTEIN C11C11.02 IN CHROMOSOME II	NCS-E10082-310500-021-F10 ET0082 Homo sapiens cDNA	NCG-E 10082-310500-021-F10 ET0082 Homo saplens cDNA	S. Occidentalis INV gene for Invertase (EC 3.2.1.26)	cds	GREEN-SENSITIVE OPSIN (GREEN CONF PHOTOBEOFERTOR BILLIAMENTE	GREEN SENSITIVE OBSIN (OBSERVOOR) TO CHECET ION PIGMEN I) (RFH-G)	Mathematical James Control (NECEPTOR PIGMENT) (KFH-G)	WON WILL FEBS. 19 Integrals Social 123 of 160 of the complete genome	YOU WILLEDIAIND FACIOR PRECURSOR (VWF) Homo sapiens PHEX gene		Homo sepiens NRAMP2 gene, for natural resistance-associated macrophage protein 2	CELL DIVISION PROTEIN FTSW HOMOLOG	Human lambda-immunoglobulin constant region complex (germline)	Mus musculus Jerky (Jrk), mRNA	601511052F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE 3012812 F	Homo sapiens Na/K-ATPase gamma subunit (FXYD2) gane complete ale alle alle	Cavia cobaya mRNA for serine/threatine kinese complete cas, alternatively spliced	HYPOTHETICAL PROTEIN MG039.	Homo sapiens ATP synthase, H+ transporting, mitochondrial F1 complex, delta subunit (ATP6D), nuclear gene encoding mitochondrial provision many a
igle Exon Prol	Top Hit Database Source	300	EST HUMAN	IN.	1	Т	SWISSPROT	Т	Т		T HUMANI	Т	T	Т	Т		Ę	SWISSPROT	SWISSPROT	Т	TOBBOT	7		╗	ISSPROT			T_HUMAN			SWISSPROT H	
.ig	Top Hit Acession No.	DE440064 4	A 420077 0	2.0E-01 AL 139077.Z	28E-04 B4028E 4	2.0E-01 IN 10363.1	R02411 1	2.6E-01 BF144331 1	XR2641 1	Τ	Τ	T	2.6E-01 BF830339 1	T	2 8E-01 X17604 1	T	2.8E-01 AF057121.1							-01 AB015355.1		01 A01/00.1	8855		-			4502296 NT
	Most Similar (Top) Hit BLAST E Value	2 GE O	2,000	2,000-0	285.04	2,611.04	2.6F-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.8F-01		2.8E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01	2.6E-01 Y10198.1	L	2.05-01	200.0	2.05-01	2.05-01	2.65-01	2.6E-01 A	2.6E-01 D88425.1	2.6E-01 P	2.5E-01
	Expression Signal	0.82	0.70	0.69	10	990	1.3	1.15	0.64	0.64	3.05	2.13	4.32	4.32	0.98		0.62	1.19	1.19	0.48	0.74	6:0	- 4	4 70	27.00	7		3.1	2.81	1.58	2.19	2.55
	ORF SEQ ID NO:	33050			33413	33465	33566	33620			34064	34140	34424	34425	35116			35626	72905					37310	+	1	1	34077	, , , , ,	+	\dagger	25684
	Exan SEQ ID NO:	19972	25110	ı	20304	20351	20444	20499	20735	20736	20926	21003	21286	21288	21942		22291	227478	2412	225/8	22738	23052	23113	24008	24111	24534	25309	24693	24BOR	08047	24230	13045
	Probe SEQ ID NO:	7289	7329	7363	7639	7687	7748	7804	8040	8040	8232	8309	8594	8294	9387	-	9639	9700	3 8	200	10090	10408	10467	11400	11511	L		L	12565	1		83
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					Sing	e Exon Prop	Single Exon Probes Expressed in brain
Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
100	13045	25684	2.39	2.5E-01	4502296 NT		Homo sapiens ATP synthase, H+ transporting, mitochondrial F1 complex, deta subunit (A 1 F o.), indicatione encoding mitochondrial protein, mRNA
248				2.5E-01	2.5E-01 M26501.1		Starfish (P.ochraceus) cytoplasmic actin gene, complete cas
2 6	l _	28250				LΝ	Mus musculus ICR/Swiss glyceraldehyde 3-phosphate dehydrogenase (Gapd-S) gene, complete cds
1038	L						Ureaplasma urealyticum section 57 of 59 of the complete genome Ureaplasma urealyticum section 57 of 59 of the complete genome
1099	L		6.42			HUMAN	ye11g07.T Stratagene lung (#557.210) nunis sapiens con oracio importation
1509		26941	6.0			LN	Boulyd's circles august 14 circles and 15 circles a
1721	14464		4.79	25年	35406	LZ L	Homo sapiens hyperpolarization activated cyclic nucleotide-gated potassium channel 4 (HCN4) mRNA
1876		27323		2.5E-		EST_HUMAN	PM4-CT0400-310700-008-008 CT0400 months septems cDNA
1876	L		1.58			EST_HUMAN	PM4-CT0400-310700-003-009 CT0400 TOTTO September CT0400 CT
2407	L	L	16			Ľ	Aquitex aedicus section 7 of 109 of ute curiprese general
2500	_		1.09			EST_HUMAN	251181271 NO COAR GOOD DOING SERVING COUNTY COME IMAGE 4282279 5
3370	1		0.84		BF698193.1	EST_HUMAN	60212552571 NIH MGC 30 name september 20NA
3407			3.04		AW973471.1	EST HUMAN	EST385404 MAGE resequences, indicator actions actions and actions and actions and actions and actions and actions and actions and actions and actions and actions and actions and actions are actions and actions and actions are actions and actions and actions are actions and actions and actions are actions as a construction and actions are actions and actions are actions and actions are actions as actions are actions and actions are actions actions are actions actions are actions actions are actions actions are actions actions actions are actions actions are actions actions actions are actions actions actions are actions actions actions actions are actions actions actions actions are actions act
3524	L	28935	1,25		AF233875.1	μ	Danio retro pepude 17 percusus 4 confine framment No 29
3537					2.5E-01 AL161517.2	Z	Arabidopsis mallana Diva chi omessinia 4, con ug inginon con con con il MAGE: 2364780 3'
3828	1_	29211			AI741483.1	EST HUMAN	Wg11c0/,XI Source INST 10 SW OT PA P ST Homo seplens cDNA clone IMAGE: 2364780 3
3828	L				ł	EST HUMAN	Wallow And Long State Comments and Comments a
4283	17022	2	0.97	2.5E		SWISSPROI	NOTE INCIDITION HORMONE PRECURSOR (MIH)
4700	L	4	1.25	2.5E		SWISSPROI	MOLT-INDIED HAVE TO WAS A Secondated protein 2 (DAP2) mRNA, complete cds
4706	1	30070	3.99	2.6	1	IN I	Viterio cholorae chromosome II. section 73 of 93 of the complete chromosome
4732	17464	4 30101		2.5E	-01 AE004416.1	Z	Mars misculus annexin V gene, intron 4 segment containing 5' LTR and gag portion of MuERV-L (murine
) if	47,403		3.7	2.5E	-01 AJZ30113.1	LN	endogenous retrovirus) element
10/4	上	30135		2.5	-01 BE896785.1	EST_HUMAN	601437468F1 NIH_MGC_72 Homo saplens CUNA clone IMAGE: 3522000 5
4/0	.L.			25.5	-01 AW873588.1	EST HUMAN	ho62f11.x1 Soares_NFIT_GBC_S1 Hamo sapiens cUNA drate in/AGE.3041337 5 3111144 W
4987				2.50	2 5E-04 S83390.1	Ā	T3 receptor-associating cofactor-1 [human, fetal liver, mRNA, 2930 nt]
5243	┙	30000			2 FE 01 A IONE345 1	Z	Homo sapiens KVLQT1 gene
5870				1	2.3E-01 A0000040.	12	Homo sepiens chromosome 21 segment HS21C007
5871	1				2.5E-01 A 1251973.1	Į.	Homo sapiens partial steerin-1 gene
652	- 1	١			TNESTAGE	Į.	Rettus norvegicus rabin 3 (RABIN3), mRNA
6945	5 19427	32442	0.79	2.55-01			

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Table 4
Single Exon Probes Everson

Single Exon Probes Expressed in Brain	Expression (Top) Hit Top Hit Acession No. Signal BLAST E No. Source Source	33011 0.88 2.6E-01 U13992.1 NT	1.29 2.5E-01 AF134110 1 NIT	33259 0.83 2.5E-01 AI 161506.2 NIT	33303 3.6 2.5E-01 AI 163282 2 NIT	33564 2.47 2.5E-01 BF109040 1	33574 0.8 2.5E-01 BE960712.1 FST HIMAN	33955 1.87 2.5E-01 BE038595 1 EST LINAM	34128 0.7 2.5E-01 P04492 SWISSDEDT	34368 3.67 2.5E-01 H53236.1 FST HIMAN	34613 0.79 2.5E-01 M88626.1 NT	35265 15.72 2.5E-01 U89651 2 NT	35266 15.72 2.5E-01 U89651 2 NIT	35253 2.06 2.5E-01/AF0RF164.1	35254 2.08 2.5E-01/A F085164 1 NIT	35858 1.39 2.5F_01 AW/FR1007 1	NAMON TO THE PROPERTY OF THE P	36315 2.13	36319 1.21 2.5E-01 X58491.1 NT	36945 3.43 2.5E-01/D50914.1 NT	1.61 2.5E-01 AF027153.1 NT	27777	37808 F 42 2 5 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	6 13 2 5E 04 14 22 24 2	1.37 2.5E.01 AL101041.2 NT	30821 137 2 FE 01 A E 170072 1	25965 1.69 2.4F-01 AA938348 4 EST 11 18484	26269 3.34 2.4E-01 BE576124.1 EST LIMANI	26700 33.63 2.4E-01/A.1289880 1 NT	26701 33.63 2.4E-01/A.)289880 1 NT	26785 1.03 2.4E-01[Y17293.1 NT	32.88 2.4E-01 AF267753.1 NT	27340 1.33 2.4E-01 AF251708.1
			<u>.</u>											35253	35254	35858		36315	36319	36945		47777	37808			30821	25965	26269	26700	26701	26785		27340
-	<u> </u>							20819	1					22081		22646	L_	┙	_[24244	24393	24490	25388	24768	25233	13323	13598	14031	14031	14111	14581	14830
	Probe SEQ ID NO:	7252	7278	7494	7536	7,44	776	8125	8296	8634	8774	9416	9416	9472	9472	8666		10441	1044	11013	164	11803	11832	11960	12365	12412	540	828	1281	1281	1363	1843	1883

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Table 4
Single Exon Probes Expressed in Brain

208170 SPLICING FACTOR, ARGININE/SERINE-RICH 4 ; contains element TAR1 TAR1 repetitive elemen h23d04.x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:3316807 3' similar to SW:PRSB_XENLA Oryza longistaminata receptor kinase-like protein, family member D, and retrofit (gag/pol) genes, complete cd 7154004.x1 NCI_CGAP_Br16 Homo sapiens cDNA clone IMAGE:3338503 3' similar to SW;SFR4_HUMAN wc62c11.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2323220 3' similar to gb:J03464 Bos taurus guanylyl cyclase-activating protein 2 (guca2) mRNA, complete cds yy55c11.rl Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:277460 5' Mus musculus DXImx48e protein (DXImx48e) mRNA, complete cds transferase, subunit II gene, complete cds; and unknown genes Tetrahymena tharmophila macronuclear gene encoding ribosomal protein L3, exons 1-2 wo33d05.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2457129 3' wo33d05.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2457129 3' Escherichia coli K-12 MG1655 section 202 of 400 of the complete genome Rattus narvegicus mRNA for alphaB arystallin-ralated protein, complete cds IMMUNOGLOBULIN A1 PROTEASE PRECURSOR (IGA1 PROTEASE Gallus gallus brain-derived neurotrophic factor (BDNF) gene, 5' end Arabidopsis thaliana DNA chromosome 4, contig fragment No. 85 PROCOLLAGEN ALPHA 2(1) CHAIN PRECURSOR (HUMAN); Drosophila melanogaster p38a MAP kinase gene, complete cds Homo sapiens HSPC142 protein (HSPC142), mRNA AV733787 cdA Homo sapiens cDNA clone cdAADE11 5' Top Hit Descriptor Hepatitis C virus genomic RNA for polyprotein, complete cds Branchiostoma floridae mRNA for calmodulin 2 (caM2 gene) Glycine max mRNA for mitotic cyclin b1-type, complete cds squifex section 12 of 109 of the complete genome 042586 26S PROTEASE REGULATORY SUBUNIT 6A; Mus musculus Wm protein (Wm) gene, complete cds Mus musculus Wm protein (Wm) gene, complete cds H.saplens AGT gene, Pstl fragment of intron 4 Bovine adenovirus 3 complete genome D.discoideum (Ax3-K) ponA gene Homo saplens serine palmitoy S.pombe swi6 gene EST_HUMAN NT HUMAN EST HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN Top Hit Database SWISSPROT HUMAN Source ISI 눋 눋 눋 눋 뉟 뉟 눋 Ż 눌 Fop Hit Acession 7661801 AF111168.2 2.4E-01 AJ133836.2 2.4E-01 BF592336.1 2.4E-01 BF002171.1 2.4E-01 AL161589.2 2.4E-01 AF030154.1 2.4E-01 AF091216.1 2.4E-01 AV733787.1 2.4E-01 AF229644.1 2.4E-01 AJ012585.1 2.4E-01 AF091216. 2.4E-01 AI698989.1 ģ 2.4E-01 AI925707.1 2.4E-01 U72726.1 2.4E-01 L43001.1 2.4E-01 X71783.1 2.4E-01 D29960. 2.4E-01 X74209. P45384 2.4E-01 2.4E-01 2.4E-01 2.4E-01 2.4E-01 2.4E-01 2.4E-01 2.4E-01 2.4E-01 (Top) Hit BLAST E 2.4E-01 Most Similar 2.16 2.46 0.73 96.0 8.16 - 4 88. 9 1.09 96.0 0.98 8.16 0.99 2.23 0.67 2.23 9.5 0.55 0.91 1.61 Expression Signal 27843 27994 27725 28213 28554 29131 30317 30865 30866 30891 31520 31627 31741 31793 32206 33004 33163 33400 34078 ORF SEQ ÖΝΩ SEQ ID 18392 14864 14894 14985 15104 15470 15494 17610 18175 18366 18366 25076 18595 18780 18830 19928 20291 2894 20081 2765 5805 5895 5999 Probe SEQ ID 2134 2185 2258 2382 2539 3129 3145 4010 4883 4883 5375 5375 5397 5569 7243 7404 7625 8247 5569 5799 6050 5597 6441

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	801877870E4 NIII 1100	SOURCE TO SET INITIALISE SO HOMO SEPTIONS CONA Clone IMAGE:4106298 5'	Camadabasha Mini McCattan	Campylobacter jejuni NCTC1108 complete genome; segment 4/6	W4450-22.X. Soares NFL_T GBC_S1 Home septens cDNA clone IMAGE:2330906 3' similar to contains	Drosophija melanmostes SKDB con consists 1	Dreamhile melanomere SKDB cons	COLLAGEN AI PHA 1/X) CHAIN PRECIPEDE	Arabidopsis theliana DNA chromosome 4 comits for month Nin- 6	Mus musculus type 1 sigma recentor dena complete ade	P. aslatica mosaic virus denomic BNA	601441421T1 NIH MGC 65 Homo septems c DNA class 184のE-2845625 21	Homo sapiens (regile 167) adda reductose (EOB) was asset of	Arabidoosis thailana ethylena-insensitive3-like4 /FII 4) mDN /	Mus musculus mRNA for putative mot protein (mot zena)	Gallus dene codino for exerting process (see a gara)	60/842848F1 NIH MGC 34 Home Splane DNA Jes IMAG Assesses III	Homo sabians chromosome 21 segment LE240084	aromatase Poephia dutata=zehra finches, oven, mBNA, 3489 - it	Mycoplasma gentralium section 35 of 51 of the complete concess	Methanococcus Jannaschii saction 138 of 150 of the complete genome	601142073F1 NIH_MGC_14 Homo saplens cDNA clone IMAGE:3808818 F1	Version of the state of the sta	Possilo nearly (mish (mish), Hinst (hinst), Hinst (hinst), and Hinst (hinst) genes, complete cds	Drassica rispus sig gene for S-todus glycoprotein, cultivar T2	Homo saplans parial Internal of the color of the color	50/175552F1 NIH MAC 17 Hours South Mac 17 Hours Sou	Human endhranoight gene complete of	Mariniabila aggroyorans dyrB gana for DNA gyrosa entrinii B	not6do6.st NCI_CGAP_Phen Home september CDNA clone IMAGE:1100843 3' similar to contains Alu	wh21b07.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:130357.3	
gle Exon Pro	Top Hit Database Source	HOT LINAANI	EST HIMAN	LN LN	FZ	NAM III PAAN		LZ	SWISSPROT	NT	NT	N	EST HUMAN	NT.	N-	LΝ	N	EST HUMAN	N	NT	トラ	LN	EST HUMAN	Į.			LN	T HUMAN			HST HIMAN	П	
ਰ 	Top Hit Acession No.	RF242794 1	2.4E-01 BF678275 1	AL 139077.2	-01 AL139077.2	-01 A1693515 1	-01 AF220067 1	-01 AF220067.1		2.4E-01 AL161494.2		01 221647.1	-	Γ		-	01 V01507.1	2	2	01 S75898.1			01 BE311893.1		1		<u></u>	Γ		2.3E-01 AB015033.1		2.3E-01 R21732.1	
	Most Similar (Top) Hit BLAST E Value	2.4F-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4€-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4E-01	2.4是-01	2.4E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3F-04 1122837.5	2.3E-01.4	2.3E-01 Y10887.2	2.3E-01	2.3E-01	2.3E-01 N	2.3E-01 A	2.3E-01	2.3E-01 R	
	Expression Signal	1.02	0.47	0.49	0.49	7.01	0.88	0.88	1.68	4.6	1.39	2.09	1.32	1.75	2.34	2.74	1.97	2.08	3.66	1.06	5	33.31	4.19	1.12	133	2.74	1.51	2.66	1.59	3.38	1.36	7.07	
	ORF SEQ ID NO:	34332		34874	1	35181	35441	35442	36202	36598	36671		37765	37801						25810		26069	2834-1		27035	27063		27903	28105	26789	28379		
	SEQ ID NO:	21189	Ш	L		22013	ı		22984	_ }		_ [24424	_	ŀ	24588	25162	25201	24992	13167	<u>2</u>	13430	2388	14305	14345	14374	14772	15166	15367	14114	15729	15847	
	SEQ ID	8487	8552	8030	9030	9463	9603	9603	10337	10687	10739	1168	11840	1391	12019	12080	12287	12400	12720	88	622	962	913	1558	1589	1628	2038	2447	2657	2827	2963	3082	

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Single Exol Flores Expressed in Start	Top Hit Descriptor	y97h10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:213283 5	GSTA5=glutathione S-transferase Yc2 subunit (5' region, intron 1) [rats, Morris hepatoma cell line, Genomic,	2212 nt segment 1 of 3]	Homo sapiens KIAA0450 gene product (KIAA0450), mKIVA	1/1/1/1/1 Soares placenta NbZHP Homo sapirans cunna cione imagra: 1450 i 7 o	Mus musculus renin (Ren-1c) gene, pramater region	Synechocystis sp. PCC6803 complete genome, 1/27, 1-1335039	Homo sapiens mitogen-activated protein kinase psedetta (Prinkin 13) ilinuira, complete cus	Homo septens nuclear transport ractor 2 (placenual protein 19) (177 19) minus	Mus musculus tulip 1 mRNA, complete ods	Homo sapiens mRNA for KIAA1512 protein, partial cds	7K30b06.x1 NCI_CGAP_Ov18 Homo sapiens cDNA clone IMAGE:3476899 3' similar to SW:GAG_SMSAV	P03330 GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P16; INNER COATTROTEIN P14; CORE.	Ofamiliaris rom1 cene	With a second second by the se	Italoma coneum smai subulit incedina nya gene	23S rRNA (Leuconostoc camosum, Genomic, 2000 nij	as27e12.x1 Barstead aorta HPLRB6 Homo capiens cDNA clone IMAGE::2318446 3' stmitar to gp:X13236 CYTOCHROME C OXIDASE POLYPEPTIDE VIC PRECURSOR (HUMAN);	as27e12.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2318446 3' similar to gb:X13238	CYTOCHROME C OXIDASE POLYPEPTIDE VIC PRECURSOR (HUMAN);	Oryctolagus cuniculus cytochrome oxtdase subunit VIa (coxVIa2) mRNA, complete cds; nuclear gene for	mitochondrial product	as42f12,x1 Barstead aorta HPLRB6 Homo sapiens cUNA cione in/AGE:2518667 5 similiar to contains Atu	repetitive element	Homo sapiens hypothetical protein FLJ20345 (FLJ20345), mrttvA	Secale cereale omega secalin gene, complete cds	Glycine max resistance protein LM17 precursor RNA, partial cds	AV719881 GLC Homo sapiens cDNA clone GLCDGB08 5	AV719681 GLC Homo sapiens cDNA clone GLCDGB08 5	Mus musculus myosin XV (Myo15), mRNA	601511573F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE;3912859 5	2a12e08.rf Soares fetal liver speen 1NFLS Homo saprens curick come invacion assessed o	Arabidopsis mailana UNA citromosomen, oung magment to to
IE EXUIT FIUNE	Top Hit Database Source	EST_HUMAN N				T_HUMAN						NT TN	2		NAMOR I STA			NT (2	EST HUMAN		EST HUMAN	Г	TN		EST_HUMAN	NT	N.	NT	EST_HUMAN /	EST_HUMAN /		П	7	LZ.
Billo	Top Hit Acessian No.	H69836.1		\$82821.1	7662133 NT	R82252.1		D90899.1	AF092535.1	1984	AB032400.1	AB040945.1			BF058381.1			\$60371.1	A1708840.1		A1708840.1		AF198089.1		2.3E-01 AI718148.1	2.3E-01 8923323	AF000227.1	2.3E-01 AF175389.1	AV719681.1	2.3E-01 AV719681.1	6754779 NT	2.3E-01 BE888071.1	2.3E-01 N80983.1	AL161558.2
	Most Similar (Top) Hit BLAST E Value	2.3E-01			2.3E-01		2.3E-01					_				Z.3E-01	_	2.3E-01	1		2.3E-01		2.3E-01		2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01
	Expression Signal	1.14		1.01	5.22	1.1	1.98	1.83	2.51	6.19	0.84	253			2.08	4.38	0.94	0.78	1 50		150		0.93		4.33	1.08	0.9	3.14	0.64	0.64	2.94			0.71
	ORF SEQ ID NO:	28780		29205		29680	l	29784		29887	L	L				31130		31374	上		31576		32330		32549	L				L	L	33278		33569
	Exon SEQ ID NO:	16122	_	16573	L	L	l	١.	١.	L	1_				١	18242	18360	١	l	⊥	18638	丄	19323	1	19522		L	ı		L	١	1	J	20446
	Probe SEQ ID NO:	2363	33	3821	3914	4316	4368	4417	4454	4517	4988	5221			5343	5443	5563	5665	7209	g	00		6558		8229	201	7188	73.15	7318	73,67	7508	7513	7652	7750

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Single Exoll Flobes Expressed in Diam	Top Hit Descriptor	Rettus norvegicus mRNA for acid gated ion channel	Pleurodeles wait distal-less like protein PWDIX-3 (PWDIX-3) IIINNA, COMPOSE COS	Rattus norvegicus mRNA for acid gated Ion channel	nec39h12.x1 Lupski_scatic_nerve Homo sapiens cDNA cione iwAvE>555550 5 sining to contain to cont	oz14a10.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1675280 3 similar to TR:Q13040 Q13040 ATP-BINDING CASSETTE PROTEIN :	Homo sapiens PPAR delta gene, promoter region	Fresh-water sponge Emf1 alpha collagen (COLF1) gene	602085608F1 NIH MGC 83 Homo sapiens clink cine in McE: 4248969 3	601462629F1 NIH MGC 67 Homo sapiens convolente invader 3868100 5	601462629F1 NIH MGC 67 Home september CONTACT	PMZ-H I 0363-281289-003-912 PL 1 0303 Homo septems CDNA	PM2-H10353-281289-003-812 F110333 FIGURE SQUEES COLOR	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 62	Xiphophorus maculatus truncated Rex1 retrotransposon reverse transcriptase (RT) pseudogene	Miss muscrifus breast/ovarian cancer susceptibility protein (BRCA1) mRNA, complete cds	Mire miserulus mixed lineage kinase 3 (Mik3) and two pore domain K+ channel subunit (Konk6) genes,	omplete eds	Mus musculus MAP kinase kinase kinase 1 (Mekry, Culipraco	Mus musculus MAP kinase kinase tinase 1 (Mettr) mrtvA, compiete cus	Human scRNA (BC200 beta) pseudogene	Human sckny (BCZOU beta) pseudogene	Human beta-c/oppasing acout (ACLIDITy) presentations of the IMAGE 648068 5	zq87c05.r1 Strategene hN I neuron (#837233) rigino saprens convocione na reconstructione	Mus musculus vinculin gene, exon 3	histamine H2-receptor (rats, Genomic, 1928 nt)	Vidua chalybeata mitochondrion, complete genome	Homo sapiens diaphanous (Urosophila, nomoreg) z (Urva 142), danouit, variant 100, mariant 100, m	Synechocysts sp. P.C. cado complete gending, 1972, 2052, 20 200000	Callus gallus 1-box containing procein (or-1501)	Osilina gaine 1 con containing process
IB EXOII FION	Top Hit Database Source		NT	NT	EST HUMAN	EST HUMAN	NT	NT	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	L Z	LZ	LZ.	E V	2	NT	NT	TN	N	N		EST HUMAN	N	NT	NT	N.	F	N I	Z
Sing	Top Hit Acession No.	AJ006519.1	U49645.1	AJ006519.1	BE475611 1	1	2.2E-01 AF187850.1	Γ	F677538.1	E618258.1		١		2 2E-01 A F020503 1	1 484582 2	2.2E-01 ALIO1302.2	100120.	J68174.1	AF155142.1	4F117340.1	4F117340.1	2.2E-01 U01307.1	J01307.1	D50604.1	AA211218.1	L13299.1	2.2E-01 S57565.1	5835974 NT	5803002 NT	01 D64000.1	01 U67087.1	01 U67087.1
	Most Similar (Top) Hit BLAST E Value	23E-01 A		2.3E-01 A		2000	2.2E-01.A	2.2年-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01 €	2.05.01	200	2.25-01/	7.45-01	2.2E-01	2.2E-01	2.25-01/	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-01	2.2E-	2.2E-4	2.2E-
	Expression Signal	235	1.76	1.55	,	1 8	3 49	252	6.3	2.02	202	4.36	4.36	1 67	10.1	/8: L	7.12	0.72	6.45	2.11	2.11		1.21				0.93		2.07			0.56
	ORF SEQ ID NO:						20032			28044			28295						29562		L		29690		30133	L			31368		31646	
	Exan SEQ ID NO:	24728	1	1	1			1 4844	15123	1	Ł	ł	1	ł	_1	┙		16848	16935	L	L	L	L	ł	1	Ł	1	Ь.	18454			18694
	Probe SEQ (D NO:	12303	12208	12403		12647	88	100	2402	2594	2594	2884	2884		2921	3387	3794	4105	4104	(32)	4232	4323	4323	4775	4774		1304 5082	5140	5659	2885	5910	591

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		_	_										_																	
Single Exon Probes Expressed in Brain	Top Hit Descriptor		num saprems gane for tukutin, complete cds	Steptococus progenes phosphatidy/gl/cerophosphate synthese (pgsA) and ABC transporter ATP-binding	Streptococcus pyogenes phosphotidy/glycerophosphate synthase (pgsA) and ABC trensporter ATP-hinding	protein (stpA) genes, complete cds, and unknown genes	Turniari giyeopriorin B gene, exon 4	Human grycophonn B gene, exon 4	Mycchesing pheumoniae M129 section 45 of 63 of the complete genome	Mus misculus mm28 M4 2000	international international promoter region	Pan Irrorbythm McOb 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	The region profile to the second seco	The mode manufacture section 25 of 136 of the complete genome	Mis miscular Ademod	MR1-TN0045-410000 006 -003 TRIDA (2)	7804fmR r1 Soarse molecular 111111111111111111111111111111111111	ARGE PROLINE BIOLID DE CHEMINE BATERINE A PROCESS CON COME IMAGE: 291591 5'	Xenomis leavis mBNA for phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like and a few phoofs like a few phoofs	Musiculus actochlock account (2007)	Brachydanio rerio ependymin here and german chain. (E-1)	CYCLIC NUCLEOTIDE GATED CHANNEL, ROD PHOTORECEPTOR, ALPHA SUBUNIT (CNG CHANNEL 3) (CNG-3) (CNG-3)	Funaria hygrometrica chloroplast-localized small heat shock protein (CPsHSP21) mRNA, complete eds:	nuclear gene ror onloroplast product	No 1009/24F1 NIH_MCC_19 Home saplens cDNA clone IMAGE:4100189 5	WRSWINS of Cumping genome	Jacobson I Outside Burst (#657/217) Homo sapiens cDNA clone IMAGE:75855 5'	Pseudomonas aeruginosa quinoprotein ethanol dehydrogenase (exaA) gene, partial cds; cytochrome c550 precursor (exaB), NAD+ dependent acetaldehyde dehydrogenase (exaA) and partial cds; cytochrome c550	synthesis A (pqqA) genes, complete eds; and pyrroloquin>	Mus musculus PHR1 (Phr1) gene, partial cds
gle Exon Prot	Top Hit Database Source	FZ	EST DIMAN			Z				L				T HI IMAM	NO.	T HIMAN	Т	Т	1			SWISSPROT		14444	NEWOL.	H IMAN	Т			
ig	Top Hit Acession No.	1 AB038490 1	2.2E-01 AV756238 1	2.2E-01 AF082738 1			T	5		Γ	Γ	Ţ.,	Ī		3247	BF376354.1	Γ		9.1	57428	M89643.1	Q90980	AF197941 1		25671	_				Ar0/1001.1
	Most Similar (Top) Hit BLAST E Value	2.2E-01	2.2E-01	2.2E-01	2 25.04	2.2F-01	2.2E-01	2.2E-01	2.2€-01 /	2.2E-01/	2.2E-01			2.2E-01	2.2E-01		2.2E-01 V		2.2E-01 A		2.2E-01 N	2.2E-01 Q	2.2E-01 A	_					2.2E-01 A	27.0-37.7
	Expression Signal	0.73	10.21	1.28	1 28	188	1.86	0.63	0.66	2.04	1.01	0.64	3.53	4.35	1.45	1.04	1.36	13.43	0.69	0.81	3.95	0.58	4.6	1.85	0.85	0.6	0.6		0.08	1.0.5
	ORF SEQ ID NO:	32381		32777	32778	32950	32951	33144	33402		33808	34279	34632		34853	34952	35024	35259	35104	35185	35200	35358	35564	35703	35941	36092	36093	90	22 120	1
	Exon SEQ ID NO:		19662	19721	19721	L	19877		l j	- 1		21141	21486	21611	21703	21786	21859	22086	21931	22017	22030	22174	22366	22503	22724	22880	22880	23018	220gg	
	Probe SEQ ID NO:	9099	6926	7029	7029	7191	7191	7386	7628	7919	7987	8449	8794	8920	9013	8608	9189	9207	8252	9263	9276	9521	9715	9853	10076	10232	10232	10268	10341	

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						011100000000	
Probe SEQ ID NO:	ဟ	ORF SEQ ID NO:	Expression Signal	Möst Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2090	L	30425	1.09	2.1E-01	-01 AE001526.1	Z	Helicobacter bylori strain 100 cardian 87 of 422 of the
5218				2.1E-01	-01 BF672695.1	EST HUMAN	602152001F1 NIH M.C. B. H. Home control of the complete gardine
6787		32559	4.1	2.1E-01	-01 AJ223392.1	LN	Doto fracilis mitro-hondria 185, ENIA conservation of the conserva
6798	19459	32481	1.92	2.1E-01	-01 U04642.1	L	Himan affaction recently 100 to 100 t
7306	19989	33065	0.65	2.1E-01	01 001958	TORGESIME	TARE CATED BOTTER III.
7306	19989	33066	0.65	2.1E-01	01 001956	SWISSPROT	VOLTAGE CATED DOTASSIUM CHANNEL PROTEIN KV3.3 (KSHIID)
7317			2.38	2.1E-01	01 AE000972 1	L	Arhamalakis filihin and 125 1120 1120
7808	20272	33380	1.94	2.1E-01	-01 AF000949.1	N	Canis familians karatin (KRT0) nana complete genome
7054		- 6					מינה מינה מינה מינה מינה מינה מינה מינה
3	20312	33425	1.22	2.1E-01	01 AF068687.1	TN	Glycine max malate dehydrogenase (Mdh-2) gene, nuclear gene encoding mitochondrial protein partial ods
7651	20316	33426	1.22	2.1E-01	01 AF068887 1	F	Chaire malate data de la company de la compa
7971	L		1.21	2 4 6 04	TIA 000000		Circuite max marte denydrogenase (Mdh-2) gene, nuclear gene encoding mitochondrial protein, partial cds
	L	†	141	2.15	Usucus/	N	Mus musculus erythrocyte protein band 4.1-like 3 (Epb4.13), mRNA
000	200	600	•				Haemophilus influenzae hmcD, putative heemodin processing protein (hmcC), putative ABC transporter (hmcB), putative heemocin structural protein (hmcA), and heemocin immunity protein (hmch) nense common
BROA		24524	1 0	Z.1E-01		\neg	spo
3	1	15045	0.86	2.1E-01/	01 AL040537.1	EST_HUMAN	DKFZp434H0614 r1 434 (synarym: https://doi.org/10.1016
8696	- 1	34532	0.86	2.1E-01		EST HUMAN	DKF2b434H0814 r1 434 (surenum: blood) Home capitals Controlled S
8857			0.45	2.1E-01 A		Т	Homo sapiens APC, dense woon 0
8936	_	34768	5.58	2.1E-01 Z35786.1			State With the control of the form of the control of the form of the control of the form of the control of the
9404		35237	0.67	2.1E-01		T HUMAN	of 1410 rd Source melananda 2Mb LM L
9404		35238	0.57	2.1E-01.N		Т	77. The state of t
9413		35262	2.31	2.1E-01 X			A thailiana mRNA for AlRanDoth models.
9518	- 1	35354	1.13	2.1E-01 A			Hamo sanjens nGSR2 mene for riban jeladilah sadjustan
10227	22875	36088	1.47	2.1E-01 2	2.1E-01 Z97067.1		Beta vulgaris mRNA for elongation factor 1-beta
102KB	2000	0770	1				DIACYLGLYCEROL KINASE, DELTA (DIGLYCERIDE KINASE) (DGK.DEI TA) (DAG KINASE DEI TA)
10284		38122	2.5	2.1E-01 P52824		SPROT	(80 KD DIACYLGLYCEROL KINASE)
77.5	1	77 00	0.87	Z.1E-01 B		THUMAN	602131427F1 NIH_MGC_81 Home saplens cDNA clone IMAGE:4270831 5
201	-		2.19	2.1E-01	8847	Į.	Homo sapiens pancreatic polypeptide 2 (PPV2). mRNA
116/2	24171	37487	1.59	2.1E-01 B		EST HUMAN	RC3-HT0822-040500-013-b/1 HT0822 Homo saniens cDNA
11870	24944		1.38	2.1E-01 X57624.1		LN FN	Drosophila melanoaster ALA-E6 DNA named region
12377	24775		2.07	2.1E-01 A	-	F	Homo seplens freque 18D oxidir reductes (EDD)
12578	25287		1.47	2.1E-01.L	L32588.1		Human granulin gene
12635	24935		1.42	2.1E-01 BI	-6-	T HUMAN	801440712F1 NIH MGC 72 Homo sentens c ONA clano MA CEl portegge pri
					1	7	CONTRACTOR OF THE PROPERTY OF

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	Top Hit Descriptor	7a59e02.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:3223034 3'	Calline mally for evena, complete ods	and services CG1.18 protein (LOC51008), mRNA	Committee of the person of the	Mus musculus Maior Histocompatibility Locus class II region	Standbook to PCC6803 complete genome, 7/27, 781449-920915	Control capiens chromosome 21 segment HS21C013	TOTILO GENERAL SERVICES OF THE	PM1-HT0422-291-299-002-c06 HT0422 Homo sapiens cDNA	Plum pox virus strain M. complete genome, isolate PS	Home rendens distribution alpha (DTNA), mRNA	Home sariens mRNA, chromosome 1 specific transcript KIAA0505	Homo septement and the symbolies mRNA, partial cds	House section R1 recentor (headyb1) gene, complete cds	Human pragrammer is a second to the complete cds; and unknown gene	Homo sapiens 1 4422 seggests gard complete genome	Medianticoccus janinascini occusi of the Sabiens cDNA clone IMAGE:3853330 5	00 (41914111111111111111111111111111111111	601449441F1 NIT INICO OF TOTAL OF THE STATE	This addition in the Hold Homo sapiens cDNA clone IMAGE:2740395 3' similar to contains element	Ap Boozal No. 2 Com 2 Co	OLD IN THE STATE OF SECURED HEAD HEAD HEAD HEAD HEAD HEAD HEAD HE	HOND saprens of noncounter and same services services	OUX END032-190500-223-603 EN0032 Homo sepiens cDNA	Homo saniens hypothetical protein ASH1 (ASH1), mRNA	HOMEOBOX PROTEIN GLABRA2 (HOMEOBOX-LEUCINE ZIPPER PROTEIN ATHB-10) (HD-ZIP	PROTEIN ATHB-10)	Ref SOD-2 gene for manganese contraining suite c	Homo saplens dual oxidase-like domains z (OCOXZ), minore	F.rubripes UNA encoding for varyet was cyningers of the complete of the comple	Saconaromyces cerevisiae i rack (1227)	Human inspancy is grown in account of the contract of the cont	
	Top Hit Database Source	CCT LIMAN 7								THE PARK	Т						- N		٦	HUMAN	Z		SSPRO		Т	TOWAR		ISSPROT	NT	NT	LN.	LN LN	7	SWISSPROI
Allio	Top Hit Acession No.	1		2.0E-01 AB017437.1	705601		-			1		-01 AJ243957.1	4503408 N	01 AB007974.1	01 AF260700.1	01 U22346.1	01 AF111170.3	2.0E-01 U67525.1	2.0E-01 BE871330.1	-01 BE871330.1	-01 X82877.1	2.0E-01 AW238005.1	P34641	2.0E-01 AL163204.2	2.0E-01 Z46906.1	BE82616	8922080 101	2.0E-01 P46607	2.0E-01 X56600.1	11432540 NT	2.0E-01 X91856.1	2.0E-01 U15300.1	2.0E-01 M75967.1	2.0E-01 P02467
	Most Similar (Top) Hit BLAST E	- 1	215-01	2.0E-01/	2.0E-01	2.0E-01 M77085.1	2.0E-01	2.0E-01 D90905.1	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	20E-01			2.0E-01						2.0E-01							
	Expression Signal		1.79	2.43	3.11	1.24	2.19	1.09	2.47	1.77	1.63	1.52	14.63	1.97	1.01	1.4	1.67	4.33	1.12	1.12	1.63	0.71	0.89				5.26	8.0				6.3		0.79
İ	ORF SEQ ID NO:		30960	25650		26103	26225	26414	26519	26651	26706		26904	26977		١			27329				29087	L			30309	28803	1	1			L	31943
	Exan SEQ ID		25019	1	13305	13458	13564	13753	13860	1	14035	14190		Ł	L	L	L	Ł	L	L	L		1	1	l	L	17702	16227	1	-1	1	Ł	_	1
	Probe SEQ ID NO:		12758	195	524	883	792	8	1103	1232	1285	143	1470	154	1550	180	1712	1752	4000	38	2347	3555	388	3822	3936	4526	4979		2	200	27.7.	2080	808	6192

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		M.auratus mu class glutathione transferase gene	PM1-CT0247-141099-001-906 CT0247 Homo saplens cDNA	Mus musculus phosphofructokinase-1 C Isozyme (Pfkc) gene, exons 3 through 7	GAMMA-GLUTAMYLTRANSPEPTIDASE PRECURSOR	Mouse germ line gene coding for beta-globin (Y2)	Andes wrus strain 0123133 glycoprotein G1 and G2 precursor, gene, partial cds	M.muscuius sopz gene exon 14	OUTS44648F1 NIH_MGC_8 Home saplens cDNA clone IMAGE:3877794 5'	Activestatum discordeum random slug cDNA19 protein (rsc19) mRNA, partial cds	Arabidopsis pyruvate decarboxylase-2 (Pdc2) gene, complete cds	Chlamydia trachomatis section 5 of 87 of the complete genome	DAUGH I FALESS PROTEIN	UAUGH I ERLESS PROTEIN	Homo sapiens filamin 2 (FLN2) mRNA, complete cds	Arabidopsis inaliana root gravitopism control protein (PIN2) gene, complete cds	Arabidopsis thallana root gravitropism control protein (PIN2) gene, complete cds	Tomo sapiens cAMP specific phosphodiesterese (PDE4C) gene, exons 2 through 12	Thomas sapiens CAMIP specific phosphodiesterase (PDE4C) gene, exons 2 through 12	D. Interanglaster UNA mobile element (hoppel)	Salvellens almana for N I K2 receptor	Salvelling plunds mkind for transferrin, complete cds	Chlorelle valued o placed -	Chlorella wildaris chloromact	Imephales promates liver afficiency and the contract of the co	Homo saniens ninein 1 m 1000 m	EST387405 MACE Percent	Ov80a10 s1 Spares testic NLT L.	Homo saniens Kirzh kirding 18 19 19 19 19 19 19 19 19 19 19 19 19 19	Mis misculus & control of the contro	Rethis negations if uclosemine 3 kinase (Fn3k), mRNA	Miss missing sold on (1) mRNA	Mono saniens lembridgings wild type alide, 3' region, partial cds.	Homo sapiens lambdelinte protein Kinase C-interacting protein mRNA, complete cds	Complete Selection of the Complete of the Comp
igle Exon Pro	Top Hit Database Source		2	EST HUMAN	TOGGGGIMIS	DAY SOLVE	- LV	Ė	í.	NAMOL - PA	LN.	F	WI COODE	SWISSPACI	SWISSPRO!	- F	- LV											T HUMAN	Т							
Sin	Top Hit Acession No.	-04 Ye4022 4	7,000,1	-01 AF250374 4	2.0E-01 P64422	2.0E-01 V00728 1	AF028028 1	2.0E-01 X91151 1	T	T]-			123	T		2.0E-01 AF157814 1	T	Γ	2.0E-01 X97121 1	T		7524759	7524759 NT	31 AF206637.2 N			Γ	Γ	8495	7649743 NT	-004353.1	Π		1
Most Similar	(Top) Hit BLAST E Value	2 OF-04	2000	20F-01	2.0E-01	20E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01/P11420	2.0E-01	2.0F-01	2.0E-011	2.0E-01	2.0E-01	2.0E-01/	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01 A	2.0E-01 A	2.0E-01 A	2.0E-01 A	2.0E-01 A	2.0E-01	1.9E-01	1.9E-01 AF004353.1	1.9E-01 U32581.2	1.9E-01 U32581.2	
	Expression Signal	3.2	402	1.28	0.68	0.84	5.8	2.95	0.99	0.82	0.68	4.97	0.65	0.65	2.11	1.98	1.98	0.68	0.68	0.69	2.78	1.56	1.56	1.4	1.4	1.61	1.39	1.36	3.58	2.68	1.87	3.9	6.86	1.43	1.43	
	ORF SEQ ID NO:		32200	32954	33102	33452		33925		35103	35129		35493	35494		36792	35793	35933	35934		36167	36674	36675	37530	37631	1		30894	30985		30978		25781	28058	28059	
<u> </u>	SEQ ID NO:	19105	19203	Ιi		1	ı				- 1				22442		1		22715	22762	22952	23431	23431	24207	24207	24762	25210	25139	24950	24924	25014	12929	13143	13420	13420	
d	SEQ ID	6335	6436	7194	7345	7878	7853	8100	8624	9251	9290	9456	9646	9846	9791	9941	9941	10067	10067	1014	10305	10744	1074		11609	12398	12546	12336	45034	12018	12763	8	342	2	641	

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Table 4
Sindle Exon Probes Expression

		_	_	-			_			_															_	_	_				_
Single Exon Probes Expressed in Brain	Top Hit Descriptor		Marsupial cat bela-dichin remain months and selections and selections and selections are selected as the selection and selections are selected as the selection and selections are selected as the selection are selected as the selection are selected as the selection are selected as the selection are selected as the selection are selected as the selection are selected as the selection are selected as the selected	Marsucial cat beta-clothin gare men.	7		7	RC5-ET0082-060700-022-402 ET0082 Home sanians CDNA	Arabidopsis thallana DNA chromosome 4 contra from a 1	Arabidopsis thaliana DNA chromosome 4 contribution 10. 10	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds. alternatively	ogetions NCI_CGAP_PNS1 Homo sapiens cDNA clone IMAGE-1637487 2	PROTHYMOSIN ALPHA (HUMAN);contains element OFR reportitive element to gb:LZ1698_cds1	0/96/02.s1 NCI_CGAP_PNS1 Homo sapiens cDNA clone IMAGE:1637487 3 similar to nhi 12489 And	The Intrincial ALPHA (HUMAN); contains element OFR repetitive element:	Raturs norvegicus sodium channel I mRNA, complete cds	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 6	imuenza A/Guangdong/243/72 nucleoprotein (seg 5) gene, 5' end	Mus musculus ATP-binding cassette 1, sub-family A, member 1 (Abca1) gene, complete ode	Urosophila melanogastar clathrin light chain mRNA, complete cds	Mus musculus p116Rip mRNA, complete cds	Mus musculus Cctg gene for chaperonin containing TCP-1 gamma subunit, partial cds. Homo sapiens calcium channel, voltage-dependent, beta 2 subunit (CACNR2) mRNA.	Products	Oryzias latipes gene for membrane guanylyl cyclase OIGC1, complete cds	Wd/110/2.x1 NC_CGAP_Lu24 Homo sapiens cDNA clone IMAGE.23370613	Urctyostelium discoldeum plasmid Ddp5, complete genome	Yersinla pestis plasmid pCD1	Mus musculus guanylate nucleotide binding protein 1 (Gbp1), mRNA	Mus musculus guanylate nucleotide binding protein 1 (Gbp1), mRNA	Homo saplens latent transforming growth factor beta binding protein 4 (LTBP4) mRNA	GAMMA BUTYROBETAINE HYDROXYI ASE.
ngle Exon Pr	Top Hit Database Source		Z	Z L		EST HUMAN	EST_HUMAN	EST_HUMAN	L	LN	LN		EST_HUMAN	FOT LIMAN	FIN FIN	I I	FIA	1	- N-	FIA		;	-	NI FOT	ES - HOMAN		Į.				EST_HUMAN
ะ ชี	Top Hit Acess No.	, 2027, 77	-01 M14568.1	1.9E-01 M14568.1	1 00 04 6 60 6	AA912486.1	1.9E-01 BE830353.1	DE630353.1	1.9E-01 AL161503.2	-01/AL161503.2	-01 AF223391.1	4 4040404	VI AMS12480.1	01 AA912480 1	T	ļ			T		-		1 8E-01 AB024400 0	l	1	4 8E-04 A1 447490 4	1	0/53947 NT	6753947 NT	I NI OSDGDG+	AI733708.1 E
	Most Similar (Top) Hit BLAST E Value		1.9E-01	1.9E-01	10 10	1.95-01	1.91-0	10-10-1	1.95-01	1.9E-01	1.9E-01	20 HO 1	1.35-01	1.9E-01	1.9E-01	1.9E-01	1 9F-01	1917-01	1.9E-01	1.8E-01	1.8E-01/	1 RF-04	1 RF-01 /	1.8E-01.0	1.8E-01.4	A BE O1	10.10	10.1	1.8F-01		1.8E-01 A
	Expression Signal	1 24	1.64	1.24	0	0.05	0.00	00.0	2:40	2.48	2.09	134		1.34	1.63	2.77	1.6	13	1.67	2.61	0.9	1.78	1.01	0.94	1,63	8.28	1 07	107	12		1.58
	ORF SEQ ID NO:	34681		34682	35632		36006	36470	38474	1	36683	36915	-	36916	37399	37645	37673	37772		25475	25700	25802	26158	26390	26485	26683	26925	26926	+		1
	SEO ID NO:	21536	ļ	21536	22426	1	22790	23237	2227		23346	23661		23661	24088	24320	24343	24431	24785	12858	16639	13159	13503	13726	13826	14015	14239	14239	14577		14097
	Probe SEQ ID NO:	8844	7700	8	9776	10142	10142	10540	10540		10655	10986		10986	11487	11726	11752	11847	12389	စ္တ	283	361	729	961	1069	1266	1492	1492	П	4050	1
																					_				_	_	_	_	_		_

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Aquarius amplus cytochrome oxidase subunit I (COI) gene, partial cds; mitochondrial gene for mitochondrial Víbrio cholerae hypoxanthine phosphoribosyfransferase (hpt) gene, partial cds, hemagglutinin/protease regulatory protein (hapR) gene, complete cds, and YRAL VIBCO gene, partial cds Bacterlophage r1t integrase, repressor protein (rro), dUTPase, holin and Iysin genes, complete cds Citrulius fanatus mRNA for wsus, complete cds P. dumerilli histone gene cluster for core histones H2A, H2B, H3 and H4 NEUROFILAMENT TRIPLET L PROTEIN (NEUROFILAMENT LIGHT POLYPEPTIDE) (NF-L) Lymantria dispar nucleopolyhedrovirus, complete genome Lymantria dispar nucleopolyhedrovirus, complete genome Homo sapiens BNIP3H (BNIP3H) gene, complete cds; nuclear gene for mitochondrial product B.taurus mRNA for potassium channel
Rattus norvegicus Thromboxane receptor (Tbxa2r), mRNA
Bovine ephemeral fever virus, complete genome
602019923F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4155318 6¹
DNA TERMINAL, PROTEIN (BELLETT PROTEIN) (PTP PROTEIN)
3/1448h10.r1 Soares placenta NbZHP Homo sapiens cDNA clone IMAGE:133027 6²
E.dispar mRNA for hexokinase (fnxt1) Scommune aradidine-5-phosphate decarboxylase (URA1) gene, complete cds
 Scommune aradidine-5-phosphate decarboxylase (URA1) gene, complete cds Rattus norvegicus CaBP9k gene 601274604F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3616788 6 Methanococcus Jannaschil section 90 of 150 of the complete genome Dictyostellum discoldeum unknown (DG1041) gene, complete cds Human carcinoembryonic antigen (CEA) gene, expn 4 S.commune orotidine-5'-phosphate decarboxylase (URA1) gene, Top Hit Descriptor M.musculus mRNA for P19-protein tyrosine phosphatase A.thaliana mRNA for ribonuclectide reductase R2 COLLAGEN ALPHA 2(1) CHAIN PRECURSOF Citrullus lanatus mRNA for wsus, complete cds Single Exon Probes Expressed in Brain AMP NUCLEOSIDASE NT SWISSPROT Top Hit Database SWISSPROT SWISSPROT EST_HUMAN HUMAN Source **EST HUMAN** SWISSPROT 눋 È Ę 눋 눋 눋 토토 Ę Top Hit Acession 8394421 10086561 1.8E-01 AB018561.1 1.8E-01 AB018561.1 ģ AF200252. BF348623.1 1.8E-01 P15272 1.8E-01 M26019.1 1.8E-01 M26018.1 1.8E-01 P08123 1.8E-01 U67548.1 1.7E-01 AF000716.1 AF255051.1 1.7E-01 AF081810. .8E-01 M59257.1 1.8E-01 X57033.1 X63440.1 1.8E-01 X77338.1 1.8E-01 X16635.1 X53330 1 1.8E-01 Q96682 P35616 1.8E-01 Most Similar (Top) Hit BLAST E 1.8E-01 1.8E-01 1.8E-01 1.8E-01 1.7E-01 1.8E-01 Value 0.94 0.91 0.75 0.77 0.78 3.08 7.28 2.61 5.69 2.64 3.45 Expression 3.28 <u>9</u> 2.29 191 1.57 2.32 1.89 Signal 35348 35349 35526 35530 ORF SEQ 36124 36465 36508 32690 36567 36870 36438 37691 ÖΝΩ 25972 26455 26456 26221 28275 SEQ ID 22126 22167 22167 22331 22335 22914 19644 22681 24358 24514 23230 19644 24908 13708 14710 15631 9514 SEQ ID 9473 9514 9679 10033 10942 9683 10633 10637 10266 10577 11439 12026 12476 10637 11767 12585 12745 787 ö 2863 988 1974

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Fig. Exp. CPF SEC CEATURE Model Similar Top Hit Accessed in Problem Top Hit Descriptor Top Hit Descriptor Top Hit Descriptor Top Hit Accessed in Problem Top Hit Descriptor Top Hit Descriptor Top Hit Accessed in Problem Top Hit Descriptor Top Hit Accessed in Problem Top Hit Accessed in Problem Top Hit Accessed in Problem Top Hit Descriptor Top Hit Descriptor Top Hit Descriptor Top Hit Descriptor Top Hit Accessed in Problem Top Hit Descriptor Top Hit Descri			_	7	7	7	7	_	-,	_	_	-	_	_	_								_	_			•	٠					•	
Delabase Continue bes Expressed in Brain			Pseudomonas putida long-chain-fatty-acid-CoA Ilgase (fadD) gene complete add	Homo saplens cleavage and polyadenylation specificity factor 3 73th submit 1000000	Homo sapiens cleavage and polyadenylation specificity factor 3, 73kill subunit (APSE2)	RC2-BN0032-120200-011-a10 BN0032 Homo sepiens cDNA	Rat (SHR strain) SX1 gene	Homo sapiens neuroligin 3 isoform gene, complete cds, afternatively splined	Homo sapiens neuroligin 3 isoform gene, complete cds, alternatively soliced	801118872F1 NIH_MGC_16 Hamo sapiens cDNA clone IMAGE:3357184 R	601116672F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3357184 5	Bacillus halodurans genomic DNA, section 2/14	ES 1389584 MAGE resequences, MAGO Homo sapiens cDNA	EST389564 MAGE resequences, MAGO Homo sapiens cDNA	Human class IV alcohol dehydrogenase (ADH7) gene, exon 3	Drosophila melanogaster mRNA for serine probasa inhibitor (seemla a) Vaca	Homo seplens chromosome 21 segment HS21C084	Homo sapiens solute carrier family 7 (cationic amino acid transporter, y+ system), member 2 (SI C7A 2)	960907.81 NC CGAP CAB Hama American Pakis	RANSFORMING PROTEIN RHOC (HUMAN).	rabidopsis thallana DNA chromosome 4, contin fragment No. 42	01286547F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3813258 F	143a03.81 NCI_CGAP_CNS1 Homo saplens cDNA clone IMAGE:1428924.3	tus musculus adenomatosis polyposis coll binding protein Eb1 (Eb1), mRNA	lus muscullus adenomatosis polyposis coli binding protein Eb1 (Eb1), mRNA	.pombe pop1+ gene	49/09.51 Soares NFL T GBC S1 Homo saplens cDNA clone IMAGE:14602973	IG RECEPTOR FORM (ARGIE SI IBLINIT BEA DELEGISEA	3G FC FRAGMENT RECEPTOR TRANSPORTER ALPHA CHAIN) (NEONATAL FC RECEPTOR)	G RECEPTOR FORN LARGE SUBUNIT PSI PRECURSOR (FCRN) (MEGNATAL EC BESERVER)	SE FC FRAGMENT RECEPTOR TRANSPORTER, ALPHA CHAIN)	will sapiens calcium channel, voltage-dependent, alpha 11 subunit (CACNA11), mRNA	And Subjects Critical Some 21 segment HS21C078	
Exon ORF SEQ Expression Crop Hill Top Hit Acess Signal D NO: Signal D NO: Signal D NO: Signal D NO:	ngle Exon Pro	Top Hit Database Source		Z.	N	N	ES I HUMAN	Ł	Z		Т	TOMAN		NAMOL .	TOMAN						T_HUMAN		J	-HOMAN			+	┰	_			DA POST		
Bear Color is	Top Hit Acession No.	, 20001714	AF 15000B.1		014/0000	D00204 4	D00364.1	AF217413.1	AE0534404	3F253142.	AP001508 4	AW/977455 1	I	Τ		T	T	11427203		7	1		0000	710000017	8391 1	1	l				11418157		1	
Exon ORF SEQ Express Signa			1 77 5	1.7E-04	17E 04	1 7E-04	1 75 04	1 75 01	1 7F-01	1 7 01	1.7F-01	1.7E-01	1.7E-01	1.7F-01	1 7E-01	1 7E-04	1011	1.75-01	1.7E-01		1.7E-01 A	4 7F 04 B	7E-01 B	1.7E-01	1.7E-01	1.7E-01/Y	1.7E-01 A	1.7E-01 P	$\overline{}$	=1-		1.7E.01		
10 SEQ OR 10 SEQ OR 11 SEQ OR 12 SEQ OR 12 SEQ OR 13 SEQ SEQ 14 SEQ SEQ 15 SEQ SEQ 15 SEQ SEQ 16 SEQ SEQ SEQ 16 SEQ SEQ SEQ 17 SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ SEQ SEQ SEQ SEQ SEQ SEQ 18 SEQ SEQ		Expression Signal	0.82	8 10	9 9	0.47	2.09	0.75	0.75	0.48	0.48	7.85	0.51	0.51	3.14	0.63	20		1.4	,	1.72	87.8	2.85	9.13	9.13	1.62	1.69	1.83	00,	1.62	1.62	2.62	1.95	
8 - : : : : : : : : : : : : : : : : : :		ORF SEQ ID NO:	L	34001	34002	34431	34462	34580	34581	34916	34917	35323	35435	35436	35455	35555	-		38154	38450	8	36511	36637	36995	36996	37098	37341		37862	33	37664	37789		
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		Probe SEQ IC NO:	786	817E	8175	8598	8628	8743	8743	808	9906	9490	9597	9597	9615	9708	10133		10293	10295	10501	10579	10709	11055	11055	11148	11348	11712	11748	1	11746	11874	12000	

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Single Exoli Plobes Light cosca in cremi	Top Hit Descriptor	tx69g05.x1 NCI_CGAP_Utf Home sapiens cDNA clone IMAGE: 22/46/23 Silling Spring	ACID RECEPTOR ALPHA-1 (HUMAN);	Human beta globin region of cition research	Homo sapiens mevalonate Kinase gene, excil o and 1		Г	Г	1	Г	Т	十	Home senior cytochrone P450 3A4 (CYP3A4) gene, promoter region	House seriors symphome P450 3A4 (CYP3A4) gene, promoter region	none square of Trichole ABI3 dene	Populus mismodal party in the Land	Populus trichocarpa cv. 1 richobel Abis yerie	Vibrio chalerae chromosome II, section 70 of 93 of the complete companies	Homo sapiens apelin gene, complete cds	T	Т	MICRONITCI FAR LINKER HISTONE POLYPROTEIN (MIC LH) [CONTAINS: LINKER HISTONE			\neg	Lycoper stoot esourchism is a second of the	Lycoperator resonant ferrain Dd2) variant-specific surface protein (var-1) gene, complete cds	Plasmodium raiciperum varam only provided in the IMAGE 2886969 3' similar to TR: 075984 075984				Rettus norvegicus CCAAT/enhancer binding brueni e-pontini (Copper) a	Г	Ī	Т	1
ie Exon Pio	Top Hit Database Source		EST_HUMAN	N	ł. I	EST_HUMAN	EST HUMAN	Ę	SWISSPROT	LIV	L'A	Į.	Z	Z	Z	눌	N.	5	L.V	MAAN LA TOT	TO TO TO THE	- N	SWISSPROT		EST HUMAN	-N	LN.	N		EST HOMAN	EST HUMAN	N.	EST HUMAN	FST HUMAN	EST HUMAN	
Sing	Top Hit Acession No.		01 AIB24404.1	01 U01317.1	01 AF217532.1	04 R31497.1	04 44548863.1	04 AE208117 1					-01 AB037729.1	E-01 AF185589.1	AF185589.1	1 6E-01 AJ003165.1	4 RE 01 A 1003165 1	AE004449 4	1.6E-01 AE004413.1	1.6E-01 AF 178000.1	1.6E-01 AW 968601.1	675331	-01 P40631		1.6E-01 AA088343.1	1.6E-01 AJ006356.1	E-01 AJ006356.1	E-01 L40608.1		1.6E-01 AW197496.1	F_01 AW 197496.1	E 04 AF034718 1	4 RE 04 RE025803 1	1.0E-01 DE402584 4	1.0E-01 DF 10304.1	ID I ISSUET:
-	Most Similar (Top) Hit T BLAST E Value		1.7E-01	1 7E-01 L	1 6E-01	4 RF-04 F	1 6F-04	1 0 30 1	1.01.0	1.05.0	1.6E-01	1.6E-01 X94Z3Z.1	1.6E-01/	1.6E-01	1.6E-01	1 6E-01	1019	1.05.01	1.6E-01	1.95-01	1.8E-01	1.6E-01	1 6F-01		-	1.6E-01	1.6	1.0				2 4			1	
- 	Expression Signal	1	65	18.27	238	2017	1.01	0 0	3.82	 8.	1.51	1.35	1.4	10.17	10.17	121	146	1.21	2.49	9.42	3.07	4.35	K 0	3	1.38	1.54		000		2.95	200					0.77
	ORF SEQ ED NO:	1		90000	30890	70007	28081	26927	26944	27364		27844	27957	L	L		01.082			29661				30218	30237			١		30947		1	١			31910
	Exen SEQ ID NO:		20,00	70107	24889	12940	15518			14654	14713	15593	L	L	L	1	- 1	16377	16730	17033	17159	L		17596	17819		1	┙	18108	5 18234	1	ı				18939
	Probe SEQ ID NO:			12268	12552	<u> </u>	8	1493	1512	1917	1977	2383	2497	7804	7000	207	3624	3624	3982	4294	4423	4431		4869	200	1007		24	5303	5435		5435	5447	5938	6162	6162

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Table 4
Single Exon Probes Expensed

		_	-,	_,							_																		•				
Single Exon Probes Expressed in Brain	Top Hit Descriptor		Arabidopsis thaliana DNA chromosome 4, contig fragment No 84	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 84	zt89d04.r1 Soares_testis_NHT Home sapiens cDNA clone IMAGE:729511 유	UI-H-BI2-agi-b-06-0-UI.s1 NCI_CGAP_Sub4 Homo sapiens cDNA close IMAGE-3724419 of	2822248.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2822248 5	AU136525 PLACE1 Homo sapiens cDNA clone PLACE1004466 6"	Gorilla gorilla androgen receptor gene, partial exon TCBAPTE0607 Pediatric pre-B cell acuts lymphobiastic leukernia Baytor-HGSC project=TCBA Homo santamo	Bacteroides vulgatus beta-lactamase (cfxA) gene, complete cds and mobilization protein (sector)	complete cds	Becillus subtilis complete genome (section 16 of 21): from 2997771 to 3213410	YOUNO8.1 Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:26a73 당	Homo sapiens guanylate cyclase activating protein (GCAP) gene exons 1-4, complete cde	S.cerevislae chromosome X reading frame ORF VJR001w	John capiana in Americani	because septemblan unnertration protein gene, partial cds; cfos gene, complete cds; and unknown gene	Scenaricio observed 189-011-h01 ST0200 Homo saplens cDNA	PM3 Litored American Kreading frame ORF YJR001w	1.0.251.1033-270100-004-11 HT0353 Homo sapiens cDNA	CUBONOCOLITING C28-G01 CT0220 Homo sepiens cDNA	CHEOMODOWAIN-HELICASE-DNA-BINDING PROTEIN 2 (CHD-2)	BO114F200E4 VIII. 1100	Description of Living Micc. 19 Homo septens cDNA clone IMAGE:3161183 5'	Fresh thought raiciparum calcium-dependent protein kinase-3 (cdpk3) gene, complete cds	inns musculus adaptor-related protein complex AP-1, beta 1 subunit (Ap151), mRNA	ouzusasesi-2 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4177073 51	Mus Intercutus protein kinase, cGMP-dependent, type II (Prkg2), mRNA	Prince of Children Sapiens CDNA clone GLCEMF07 6'	Test out item tasks Too minkN, 5 end	Circimis addis. 75. PBM: 7.0074 Homo saplens cDNA	Home controls to mixing for ent-kaurene synthase, complete cds	control capitate in this first of FLU0104 protein, partial ods
gle Exon Pro	Top Hit Database Source		LN L	LN-	EST HUMAN	EST HUMAN	EST HUMAN	ES HOMAN	EST HIMAN	NUMBER OF THE PROPERTY OF THE		Т	HOMAN		Z	Z	T CI INVANI	T	T HIMANI	Т	7	T.	T			T LI INTONI	NEW	T HI MAN		T HI MANN	- ICINOIL		
Sin	Top Hit Acest No.	0 4045000	-01 AL161588.2	1.0E-01 AL161588.2	1.0E-01 P4388047.1	T	T	T	12	T			T	I		01 AF111167.2	Ī	T	_	T,	Τ		T		1.6E-01 6871552 NT		9466	AV719585.1	T	-	T		7
	Most Similar (Top) Hit BLAST E Value	4 80 04	1.01-0-1	200.7	10.10 10.10 10.10 10.10	1919	18 19 19 19	1.6F-01	1.6E-01	1 AE-01		161-01	1 BE-01	1 6F-01		1.6E-01	1.6E-01	1.8E-01	1.6E-01	1.6E-01/	1.6E-01	1.6E-01 014847	1.6E-01 E	1.6E-01 A	1.6E-01	1.6E-011B	1.8E.01		1.6E-01 L	1.6E-01 A		1.6E-01 A	
	Expression Signal	237	2 3 2	0.55	532	1,66	0.74	1.81	0.51	0.87	0.88	0.63	0.59	172		0.83	2.09	1.7	76.0	3.3	1.59	1.59	1.55	4.28	7.28	1.26	1.64	5.28	1.72	1.5	287.76	2.4	
	ORF SEQ ID NO:	32092	L		30639	33453		33589		33844	34364	34561		34706	-		_	35391		36486	36845	36846	36852		37296			37273	31052				
	SEQ ID	19104	l	1	17944	i.	L	20464	20619	20713	21222	L	21523	21561		21699	22204	22207	22242	23249	23598	23598	23603	23729	23994	24301	25331	24538	24721	24740	25149	24901	
	Probe SEQ ID NO:	6334	6334	6685	6867	7676	7703	7768	7924	8018	8530	8725	8831	8870		RODA	9221	9554	888	10553	10918	10918	10923	11059	11388	11706	11886	12002	12292	12321	12418	12574	
																								_		_			_	_			

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t Top Hit Acession Source	01 AF28734	01 9506522 NT	04 BF672698.1 EST HUMAN	04 BE710087.1 EST HUMAN	04 BE/1008/.1 EST CIMAN	01 AV/11696.1 ESI TOWNEY	01 AL 103204.2	01 AJ009/33:1	01 AJ251885.1	01 L36125.1	01 AW195516.1 EST HUMAN	LN.		LN.	EST HUMAN	EST HUMAN	EST_HUMAN	1.5E-01 078687 SWISSPROT INADH-UBIQUINONE OXIDOREDUCI ASE CHAIR 4 HAGE-1571337 3' similar to qb:M11433	EST_HUMAN	-01 Z23104.1 NT	1.5E-01 [Z23104.1] L. Stagnalis mrtNA for G protein-coupied receptor	-01 U09964.1 NT	-01 7108358 NT	EST HUMAN	UT AWGGGGG.	-01 AW366859.1 EST TOWAR	- 2	TEGI HOMAN	E-01 BF695381.1 EST HUMAIN	E-01 Z72608.1
	1 AF287344.		1 BF672698.	1 BE710087	1 BE /1008/	1 AV/11690	1 AL 103204	1 AJ009/35	1 AJ251885	1 136125.1	1 AW19551	1 D26535.1	1 D28535.1	1 AF117340	1 AW 44445	1 BF695381	AW67251	1 078687	01 AA93504	1 ZZ3104.1	31 Z23104.1	01 009964.1		A A VAI A RES	OT AWGGGS	01 AW3868	01 AL16328	01 BF68766	01 BF69538	01 Z72608.1
Most Similar (Top) Hit BLAST E Value	1.6E-0	1,6E-0	1.85.0	1.85.0	1.58.0	1.5F.	7 S	1.5F-0	1.58.0	1.88-0	1.5E-0	1.5E-0	1.55.0	1.5E-0	1.5E-0							1.56	1.5		<u>.</u>	- -				5.
Expression Signal	1.72	1.7	1.52	1.4	1.4	9.31	1.09	1.01	2.75	1.42	0.82	2.96	2.96	1.86	1	1.98	1.15		80.8			2.11								1.55
ORF SEQ ID NO:		30892		25689	25690		26198	26487	26491		26610	26668	26669	26901		28162		28459		28777	1	<u> </u>							28162	30467
SEO S OS OS OS OS	24961	24973	25046	13050	13050	ll		. !	l	ı	L	1_	L	L	1_	15423	Į.	1	١	16120	1	l		l	16599	3 16773	16901	3 17410	3 15423	
Probe SEQ ID NO:	12881	12687	12796	241	241	573	768	1070	1075	1091	1194	1252	1252	1485	1901	2716	Ş	3048		3347	3381	9779		3752	3848	4028	4181	4676	4703	5132

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Influenza B virus (B/Nanchang/480/94) NB protein gene, complete cds; and neuraminidase gene, partial cds SEX HORMONE-BINDING GLOBULIN PRECURSOR (SHBG) (SEX STEROID-BINDING PROTEIN) (SBP) (TESTIS-SPECIFIC ANDROGEN-BINDING PROTEIN) (ABP) ob73f02.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1337019 3' similar to contains element ULHF-BNO-akk-4-05-0-ULTNIH MGC 50 Homo sapiens cDNA clone IMAGE:3077409 5'
ULHF-BNO-akk-4-05-0-ULTNIH MGC 50 Homo sapiens cDNA clone IMAGE:3077409 5'
ULHF-BNO-akk-4-05-0-ULTNIH MGC 50 Homo sapiens cDNA clone IMAGE:3077409 5'
Saccharomyces cerevisiae weak multicopy suppressor of lost-1 (SOL3) gene, complete cds
MEROZOITE RECEPTOR PK69 PRECURSOR (68 KD PROTECTIVE MINOR SURFACE ANTIGEN) Archaeoglobus fulgidus section 88 of 172 of the complete genome
Homo sepiens chromosome 5 open reading frame 3 (C5ORF3), mRNA
GLUTAMATE—CYSTEINE LIGASE REGULATORY SUBUNIT (GAMMA-GLUTAMYLCYSTEINE
SYNTHETASE) (GAMMA-ECS) (GCS LIGHT CHAIN) nw30d10.s1 NCI_CGAP_GCB0 Homo sepiens cDNA clone IMAGE:17241971 3' HYPOTHETICAL 61.7 KD PROTEIN IN THRC-TALB INTERGENIC REGION (ORFS) Mus musculus transforming growth factor alpha (TGFa) mRNA, complete cds
Mus musculus transforming growth factor alpha (TGFa) mRNA, complete cds
Mus musculus DNA methyltransferase 2 (Dnmt2), mRNA
Mus musculus DNA methyltransferase 2 (Dnmt2), mRNA
Mus musculus genomic fragment, 279 Kb, chromosome 7
601684322F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3833981 6'
Homo saplens RAD54 (S.cerevisiae)-like (RAD541) mRNA Bos taurus Niemann-Pick type C1 disease protein (NPC1) mRNA, complete cds Bos taurus Niemann-Pick type C1 disease protein (NPC1) mRNA, complete cds Morone saxatilis gonadotropin-releasing hormone type II gene, complete cds wr52c08.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2491310 3 Caiman crocodilus MHC class II beta chain (hellbeta) gene, complete cds Top Hit Descriptor EST382376 MAGE resequences, MAGK Homo sapiens cDNA Homo sapiens HARP (HARP) gene, exon 17 and complete cds L3-C70219-160200-064-F10 CT0219 Homo saplens cDN THROMBOSPONDIN 1 PRECURSOR Single Exon Probes Expressed in Brain LTR2 repetitive element Top Hit Database Source EST_HUMAN EST_HUMAN SWISSPROT SWISSPROT EST_HUMAN SWISSPROT EST_HUMAN SWISSPROT EST_HUMAN **SWISSPROT** EST_HUMAN EST HUMAN HUMAN SWISSPROT 눌 Ż 4506396 NT ξ 11417236 NT Top Hit Acession 6753650 AW850754.1 1.5E-01 P07996 1.5E-01 AF256652.1 BE727658.1 AE001039.1 1.5E-01 AF210842.1 1.6E-01 AI973157.1 1.6E-01 AF299073.1 1.5E-01 AF299073.1 4,1276505.1 AF134907.1 ġ AW970295.1 1.5E-01 AW 500611.1 1.5E-01 AW 500611.1 1.5E-01 U65016.1 1.5E-01 U65016.1 1.5E-01 AA811545.1 1.5E-01 Q28462 1.5E-01 AA714760. P15196 .6E-01 U46560.1 1.5E-01 P48508 P30143 1.5E-01 (Top) Hit BLAST E Value 1.5E-01 .5E-01 1.5E-01 1.6E-01 1.5E-01 1.6E-01 1.6E-01 Aost Simila 1.5E-01 1.6E-01 1.6E-01 1.6E-01 2.16 6.92 3.09 1.93 8.42 1.95 2.21 2.35 1.25 6.82 6.82 Expression 2.07 2.98 2.04 2.04 1.04 1.04 96.0 Signal ORF SEQ ID NO: 30480 30499 30632 31189 31653 31654 31693 31999 32201 31847 32214 32260 32361 32389 33262 33263 30554 33254 33414 33775 SEQ ID 17984 18011 18291 18734 18880 19025 25089 18700 18700 18929 19205 19216 19348 20162 19269 19376 25102 19973 20162 8 Probe SEQ ID NO: 5145 5203 5915 5915 5962 6102 5451 6492 6152 6261 6409 8448 8493 6685 6812 6882 6918 6437 7115 7480 7499 7640 490

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Single Exon Probes Expressed in brain	Top Hit Acession Database Source Source		3.1 EST HUMAN	ESI HUMAN	Z	TN	SWISSING.		EST HUMAN	7	EST_HUMAN	EST_HUMAN	N	Ŀ	L.	1	L.		401 AI814046.1 EST_HUMAIN GALGTOSIDASE-RELATED PROTEIN PRECURSOR (HUMAN); GALGTOSIDASE-RELATED PROTEIN PRECURSOR (HUMAN); GALGTOSIDASE-RELATED PROTEIN PRECURSOR (HUMAN);	ON A1814046.1 EST HUMAN GALACTOSIDASE-RELATED PROTEIN PRECURSOR (HUMAN);	۲N	NT		EST_HUMAN	EST_HUMAN	<u>L</u>		N	EST THUMAN TOWNS TOWN TO THE STATE OF THE ST	1.5E-01 AA425488.1 EST_HUMAN contains element MER22 repetitive element;
	Top Hit	4A9703	BE8847	C16800	27835.	D84476	01 P43446		N74226	BF5854	AV7548	AU1300	U0045£	į	M//14	ALOUA	AF007.	7886X	AI8140	AI8140	U4093	AJ011	AJ011	BEOBB	BE088	AL163	AL163	AB04;	AW84	1 AA 42!
	Most Similar (Top) Hit BLAST E Value	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.51.01	1.21-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01	1.5E-01		1.5E-01	1.95-01	1.5	1.5	1.5E	16	1.5					L	1.6	1.5	1.5g	
	Expression Signal	1.13	0.95	11.54	1.6	1.65	0.71	1.16	2.88	-	2.3	0.74	7.32		0.53	7.51	7.51	2.92	2.16	2 18	201	1.43				7.31		1.7	1.6	1.95
	ORF SEQ ID NO:	33947			34162	34325		34575	34842	34945			33543			35674		35965	36070	L_		L							36944	36999
	SEQ ID	20812	20905	20993	21025	21183	21204	21429	21692	21781	21788	21972	20424	13103	22368	22472	22472	22751	22855		00077	L	1	_L	L	L	L	1_		1 1
	Probe SEQ ID NO:	2 2	8211	8289	8332	268	8512	8737	0000	1000	6	9305	2000	323	9717	9821	9821	10103	40207		10207	10203	920	1050	10505	10728	10726	10896	11012	11057

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Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, qe72e01.x1 Soares_fetal_Jung_NbHL19W Homo sapiens cDNA clone IMAGE:1744536 3' similar to Candida tropicalis DNA for mitochondrial NADP-linked isocitrate dehydrogenase, complete cds Candida tropicalis DNA for mitochondrial NADP-linked isocitrate dehydrogenase, complete cds UI-H-BI1-act-a-09-0-UI.s1 NCI_CGAP_Sub3 Homo saplens cDNA clone IMAGE:2714009 yp87e04.r1 Scares fetal liver spleen 1NFLS Homo sapiens oDNA clone IMAGE:194430 6 vd54c01.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:112032 3 hr67c02.X1 NCI_CGAP_KId11 Homo sapiens cDNA clone IMAGE:3133538 3'
AU117147 HEMBA1 Homo sapiens cDNA clone HEMBA1000769 5'
AU117147 HEMBA1 Homo sapiens cDNA clone HEMBA1000769 5'
x071d12.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2581751 3' Thermotoga maritima section 22 of 136 of the complete genome Mus musculus transcription factor USF2 (USF2) gene, exons 8-10 and complete cds ye15c11.s1 Stratagene lung (#837210) Homo septiens cDNA clone IMAGE:117812.3 Sus scrofa mRNA for socium lodide symporter
Homo septens T cell receptor beta locus, TCRBV8S5P to TCRBV21S2A2 region
Xenopus laevis mRNA for DNA (cytosine-5-)-methytransferase, complete cds yg97a03.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:41467 6 yg97a03.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:41467 bx56c02.r1 NCL_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2273570 3' bx56c02.r1 NCL_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2273570 3' ny72d07.s1 NCL_CGAP_GCB1 Home saplens cDNA clone IMAGE:1283821.3 ge:M17887 60S ACIDIC RIBOSOMAL PROTEIN P2 (HUMAN); 602128753F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:4286549 5' Rattus norvegicus chemokine CX3C mRNA, complete cds GLYCEROL 3-PHOSPHATE ACYLTRANSFERASE PRECURSOR (GPAT) wm74d01.x1 NCI_CGAP_UIZ Homo septens cDNA clone IMAGE;2441665 3 w 62c08x1 NC_CGAP_Ut1 Home sepiens cDNA clone IMAGE:2491310 3 Campylobacter Jejuni NCTC11168 complete genome; segment 1/6 Top Hit Descriptor Thermotoga maritima section 22 of 136 of the complete genome Mus musculus mRNA for death inducer-obliterator-1 (Dio-1 Mus musculus growth differentiation factor 5 (Gdf5), mRNA AV741272 CB Homo saplens cDNA clone CBDAGD04 5 Single Exon Probes Expressed in Brain complete cds) EST_HUMAN HUMAN Top Hit Database Source EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN SWISSPRO-EST HUMAN EST 눋 Ż EST **Top Hit Acession** 6679980 AF030358.2 AJ238332.1 AB026898.1 Š 1.5E-01 AL139074.2 AF009663.1 AV741272.1 .5E-01 AI193704.1 1.4E-01 AE001710.1 AA720615.1 1.4E-01 AW082796.1 .4E-01 AI699094.1 1.4E-01 D78638.1 1.4E-01 AI933496 4E-01 A 699094 1.4E-01 AB004556 **U12283.1** 1.5E-01 1.5E-01/ (Top) Hit BLAST E 1.55-01 1.5E-01 1.5E-01 .4E-01 .4E-01 Most Simila .4E-01 4.01 1.4E-01 1.4E-01 1.4E-01 .4E-01 .4E-01 1.4E-01 Value 1.37 1.56 6.35 9.97 9.2 1.89 3.62 2.48 1.38 Expression 9 4.23 0.96 0.96 8.69 8.69 5.48 9.9 5.6 5.6 33051 ORF SEQ ID NO: 30965 29268 30342 30656 30679 30680 32161 32244 SEQ ID 19973 24222 25202 24739 24743 25220 13098 **1** 14635 15196 13659 14487 ö 16953 16895 18030 19244 SEQ ID 11625 12369 292 8 1745 2573 898 1978 2478 4153 4212 4153 ë 5223 5246 5246 6205 6477 6391 6391

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Probe SEQ ID NO: NO: NO: 6490 6609 77284 7521 7653 8373 8881 8881 8882 9021 9025 9246 9246 9246 9246 9258 9350 9704	Exon SEQ ID NO: 19257 19257 19274 198718 20317 2 20192 2 21550 2 21774 2 21774 2 21774 2 21875 8 2 21825 6	ORF SEQ ID NO: 32275 33283 33427 34650 34650 34654 34664 34664 34665 35096 35096 35096 35096 35096 35096 35096	Expression Signal 1.84 2.45 0.65 0.65 0.65 0.57 0.59 0.57 0.59 0.59 0.59 0.59 0.59 0.59 0.59 0.59	Most 8 (10p BLA Van Van Van Van Van Van Van Van Van Van	Similar Top Hit Acession ST E No. SING Sine Top Hit Acession No. ST E No. SING ST E No. SING ST E No. SING ST E No. SING ST E No. SING ST E No. SING ST E No. SING ST E NO	Source Source Source Source Source Source Source EST HUMAN EST HUM	Top Hit Descriptor Top Hit Descriptor Top Hit Descriptor GV1-UM0036-080300-103-d09 UM0036 Homo sapiens cDNA GV1-UM0036-080300-103-d09 UM0036 Homo sapiens cDNA GV1-UM0036-080300-103-d09 UM0036 Homo sapiens cDNA GV1-UM0036-080300-103-d09 UM0036 Homo sapiens cDNA GV1-UM0036-080300-103-d09 UM0036 Homo sapiens cDNA clone IMAGE:2710289 3' GV1-UM0036-0-ULI-s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:1879683 3' GV3-UM0036-0-ULI-s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:1879683 3' GV3-UM00371 Coares_NEF F8 gW_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:1878683 5' GV3-UM00371 COA-COA-COA-COA-COA-COA-COA-COA-COA-COA-
9887	7 22537	35732			1.4E-01 AF023813.1	TN	Macing Maritian Control Cochles Homo Sapiens cDNA clone IMAGE:2485094 5'
9704					X66092.1	F2	C.perfringens ORF for putative memorane transport protein. Macromitrum levatum small ribosomal protein 4 (rps4) gene, chloroplast gene encoding chloroplast protein.
8					AF023813.1	Ŋ	partial cds partial cds
8868					1.4E-01 AW021908.1	EST HUMAN	dr28/108.y1 Warton Fetal Cochlea Home sapiens cDNA clone (MAGE: 2485094 5
10157	22636	35847			1.4E-01 BF375285.1	EST HUMAN	MR3-ST0218-211289-013-a08 ST0218 Homo sapiens dDNA
5 5		l		Ц	1.4E-01 BF375285.1	EST HUMAN	MR3-S10218-211289-013-806 S10216 Holling September CONT. 472403 11 Scrapes fetal liver soleen 1NFLS Homo sapiens cDNA clone IMAGE:111365 51
10360	L				1.4E-01 T84293.1	EST HUMAN	yd4/dus.i I States tetal into Specific 14 of 21); from 2599451 to 2812870
10489		45 36372			1.4E-01 Z99117.1		Bacilius subulire cumprete general (CCS) Mono sapiens cDNA clone IMAGE:13203643'
10607	L				1.4E-01 AA811480.1	EST HUMAN	TANABEL 1 Spares breast 2NbHBst Homo sapiens cDNA clone IMAGE:154088 5
10746	L				1.4E-01 R53400.1	EST HUMAN	W/Jocusi I States Diess Dan 1201 Pario sapiens cDNA clone IMAGE:2803274 3
10954	54 23631	31 36879	1.31		1.4E-01 AW104982.1	EST_HUMAN	XXI OB LOCAL COLL COLL COLL COLL COLL COLL COL

Page 106 of 536 Table 4 Single Exon Probes Expressed is

Single Exon Pricession Top Hit Acession Top H				Т	T 1.	_	_						_	_	_		_				_	_									
SEQ ID ORF SEQ Expression Top Hit Acession	bes Expressed in Brain	Top Hit Descriptor	7 d Comme for 11.	Vert 910 1 State Street liver spiech TNFLS Homo sapiens cDNA clone IMAGE:120830 6'	INTEGRIN ALPHAS PRECURSOR (FIBRONECTIN RECEPTOR ALPHA SUBUNIT) (INTEGRIN ALPHA- FIVAL AS A CALLARY OF THE CONTROLL OF THE SUBUNITY (INTEGRIN ALPHA-	(r)(vLA-0)(CD48E)	C. beringens O'K for butative membrane transport protein UI-H-BiD-sal-c-08-0-11 of MOT COAR S. 14 11	Borrella burgdorfert glyceraldehyde-3-phosphate dehydrogenase (GAPDH), phosphoglycerate kinase (PGK).	M. musculus 016K dana for 18 P.D. scottle.	ביי פיינים ביינים Mus musculus neuronedin U precursor (Nmu) gene, partial cds; tPhLP (Tphip) gene, partial cds; CLOCK	Notice of the second of the se	144 AND AND COAR SIDE SEPTION CONTRACTOR (MAGE: 2972319 3)	Third the Code Code Code Code Code Code Code IMAGE: 2972319 3	Sealing Models and control additions to the control of the control	1 799 aloh nama ining	MANAGERABEA NILL MOS & U	Togu rubripes putative neurotransmitter receptore, YDR140w homolog, and dischamilde riboning services.	ransformylase (GART) genes, complete cds	29/18/2002/8/18 sp. PCC6803 complete genome, 23/27, 2868767-3002966	AND THIS THIS TO BE IN THANKS I HANSFORMING PROTEIN ABL	Ald Hildscules History for prolicese, complete cds	Igno sablens G brotein country and a country of the	iomo capidano O protein coupidad i aceptor 30 (grand) MRNA	Ome septicals of protein-coupled receptor 50 (GPR60) mRNA	iono seprens gene for NBS1, complete cds	unitari carcorrus HU/NLV/Girlington/83/UK RNA for capsid protein (ORF2), strain HU/NLV/Girlington/83/UK	uman oalicivirus HUNLV/Griington/93/UK RNA for capsid protein (ORF2) strain HUNLV/Griington/93/UK RNA for capsid protein (ORF2)	dunerill histone gene duster for one histones H2A H2R H3 and H4.	Raftus norvegicus A-kinase anchor protein mBNA complete ade	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation	
Exon Signal NO: Signal Si	gle Exon Pro		FST HIMAN	EST HIMAN	TOGGGGIMA	PALEGIMO	EST HUMAN	l F2	L			HUMAN	Т	A CANONIA			HIMAN			T	T	Г									
Exon No: ORF SEQ Expression (Top) H Nost Sm No: Most Sm No: Alue No: Signal No: Most Sm No: Alue No: <	us	Top Hit Acession No.	T96102.1	T96102.1	PORBAR		T					T	T	T	T	1968117		Γ				Γ	8467	4758467 N			T	=			AL117078.1 NT
Exon Sec ID ID NO: Signed Signed NO: Signed Signed NO: Signed Signed NO: Signed		Most Similar (Top) Hit BLAST E Value	1.4E-01	1.46-01	1.4E-01	1.4E-01	1.4E-01	1.4E-01	1.4E-01		1.4E-01/	1.4E-01	1.4E-01	1.4E-01 A	1.4E-01	1.4E-01	1.4E-01 B	200	4 4F 64 D	1.4E-01 P	1.4E-01 D	1.4E-01 A	1.3E-01	1.3E-01	1.3E-01 A	1.3F-01 A					1.3E-01 AL
Exon OR SEQ ID III SEQ III SEQ ID III SEQ II		Expression Signal	1.3	1.3	2.36	1.66	1.41	2.37	1.82		1.83	1.31	1.31	1.98	2.03	2.2	2.84	1 25	2.87	3.15	6.28	2.37	3.12	3.12	28	1.05		1.05	0.92	1.8	1.34
		ORF SEQ ID NO:				L					37661	37747	37748	30904	31049	-					-		25758	26767	25932	26035	-	20036	40707	26313	26425
Probe SEQ ID NO: 11038 11038 111038 111038 111448 111627 11627 11627 11627 11627 11627 11627 11627 11627 11627 11627 11627 11627 11627			L				19967		24112	_	24335	24411	24411	26172	24708	24714	25383	24794	24801	25407	25221	25033	13118	13118	13300	13400	- 6	13400	1000	3043	13/60
		Probe SEQ ID NO:	11036	11036	11038	11262	11301	11446	11612		11743	11827	11827	12213	12261	12275	12318	12413	12426	12500	12708	12779	314	344	516	621	200	708	200	4/8	cont

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1105	13862		2.6	1.3E-01	AL115265.1	NT	Botrytis cherea strain T4 cDNA library under conditions of nitrogen deprivation
1193	13945	5090	1.13	1.3E-01	AV712467.1	EST_HUMAN	AV712467 DCA Hamo saplens cDNA clone DCAAFF05 5'
1425	14172		1.18	1.3E-01	AF146277.1	LΝ	Homo sapiens adapter protein CMS mRNA, complete cds
1850	14588		76.0		6680957 NT	LN	Mus musculus procollagen, type XI_alpha 1 (Cd11a1), mRNA
1952	14687	27400	2.18	1.3E-01	AL117078.1	TN	Botrytis cinerea strain T4 cDNA library under conditions of nitrogen deprivation
					, OE207 01 4	1	Rhodopseudomonas acidophila pucB5, pucA5, pucB6, pucA6, pucB7, pucA7, pucB8, pucA8 and pucC
2167	Ш		1.22	1.35-01		Z	genes and OKF101
2288			1.2	1.3E-01	AW812104.1	EST_HUMAN	RC4-S10173-191099-032-d12 S10173 Homo sapiens cDNA
. 2379			3.34		AE001016.1	NT	Archaeoglobus fulgidus section 91 of 172 of the complete genome
2592			4.76	1.3E-01	M86918.1	NT	Carassius auratus keratin type I mRNA, complete cds
3065	15831	28474	1.01		AL163207.2	IN	Homo saplens chromosome 21 segment HS21C007
3443	16139	28849	86'0	1.3E-01		TN	Bovine branched chain alpha-keto add dihydrolipoyl transacylase mRNA, complete cds
3969	16718		1.43	1.3E-01	AL161581.2	IN	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 77
4117	16859		1.27		AF020713.1	IN	Bacteriophage SPBc2 complete genome
4137	16879		4.24		AW364341.1	EST_HUMAN	QV3-DT0018-081299-036-a03 DT0018 Homo saplens cDNA
4145	16887	29518	2.03	1.3E-01	AF026805.1	ΙN	Schistosoma mansoni fructose bisphosphate aldolase mRNA, complete cds
4163	16903	29532	18.52	1.3E-01	AW273741.1	EST_HUMAN	xx23f10.x1 Scares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2813995 3'
4257	16998	29627		1.3E-01	AV752279.1	EST_HUMAN	AV752279 NPD Homo sapiens cDNA clone NPDAZE02 5'
4257	16998	29628		1.3E-01	AV752279.1	EST_HUMAN	AV752279 NPD Homo sapiens cDNA clone NPDAZE02 5'
4279	L		12.76	1.3E-01	AL163280.2	ĮN.	Homo sapiens chromosome 21 segment HS21C080
4445	17181	29806	77.0	1.3E-01	M21572.1	IN	Bovine branched chain alpha-keto acid dihydrolipoyl transacylase mRNA, complete cds
4497	17233	29863	2.68	1.3E-01	BE272339.1	EST_HUMAN	601126096F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:2890063 5'
4963	17688		0.74	1.3E-01	BF091980.1	EST_HUMAN	RC4-TN0077-180900-012-c05 TN0077 Homo saplens cDNA
	·						ha07b06.x1 NCI_CGAP_Kid12 Homo sapiens cDNA done IMAGE:2872979 3' similar to contains L1.b1 L1
5242	18048	30677	0.83	1.3E-01	AW466988.1	EST_HUMAN	L1 repetitive element;
5278	18083	30739	2.23	1.3E-01	AW804417.1	EST_HUMAN	QV0-UM0093-100400-189-a06 UM0093 Homo capiens cDNA
5414	18213		0.77		AF107793.1	NT	Emericella nidulans DNA-dependent RNA polymerase II RPB140 (RPB2) gene, partial cds
5497	18295		0.75	1.35-01	AF056880.1	LN	Hepatitis C virus 68_CL10 genome polyprotein gene, partial cds
5638	18433	31346	26.0		BF210920.1	EST_HUMAN	601874591F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4101119 5'
5896		31628			_	EST HUMAN	602039337F2 NCI_CGAP_Brn67 Homo sapiens cDNA done IMAGE:4177233 5'
5896		31629			BF527281.1	EST_HUMAN	602039337F2 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4177233 5'
6392		32162	l ,		AB031326.1	۲	Schizosaccharomyces pombe gene for Alp41, complete cds
6474	19241	32241	1.95		X88891.1	LN	Cjacchus Intron 4 of visual pigment gene (red allele)
6691	19608		0.75	1.3E-01	W26367.1	EST_HUMAN	26f3 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA

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						gle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6914	19651		0.99	1.3E-01	1.3E-01 BF529560.1	EST HUMAN	602044345F1 NCI CGAP Bm87 Home servious CDNA class MACE 418,000 mi
7162			1.96	1.3E-01	-01 H48664.1	EST HUMAN	1/33402.r.1 Scares felal liver scheen 1NFI S Home scales on ANA shape 1/4/07-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2
7859	20554		0.88	1.3E-01	-01 BE272339.1	EST HUMAN	601126096F1 NIH MGC 9 Home saniers china septemble control invitors in the control of the contro
7873			1.34	1.3E-01	11423294 NT	NT TN	Homo saplens PRO0611 ordein (PRO0811) mRNA
7902		33727	1.17	1.35-01	-01 BF690522.1	EST HUMAN	602187015T1 NIH MGC 49 Home septems cDNA clone IMAGE:4280024 2
8136	. 1		0.51	1.3E-01	-01 BE562528.1	EST HUMAN	601335829F1 NIH MGC 44 Homo seniens cDNA clone IMAGE:388097 5
8172		33998	0.64	1.3E-01	1556	L	Homo saplens TED protein (TED), mRNA
8243			4.47	1.35-01	-01 Z74102.1	Z	Scerevisiae chromosome IV reading frame ORF YOU 054c
8285			4.44	1.3E-01	8923919	LN	Homo saplens core histone macroH2A2 2 (MACROH2A2) mpNA
8426	21119	34258	1.02	1.3E-01	01 BF690522.1	EST HUMAN	602187015T1 NIH MGC 49 Homo septiens cDNA clone IMAGE-43000743
8847	21538	34683	0.58	1.3E-01	01/R11172.1	EST HUMAN	y39g11.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:129284 5' similar to SP-RI 28 RAT P25334 คณร RIROSOMAN เป็นกระเมา
8847	21538	34684	0.58	1.35-01		EST HUMAN	1/39911.r1 Sogres fetal liver spleen 1NEA Homo saplens cDNA clone IMAGE:129284 5' similar to SP-RI 28 RAT Post-8 ROS DIRACEANAL DECITION
9119	1	34973	0.61		11068003	L	Plutella wiostella granulovinis, complete denome
9119	21807	34974	0.61			N-	Pittella xytostella cranuloutrus complete generale
9372	21947	35120	3.71	1.3E-01	01 AF023129.1	Į	Oroctolegius cuniculus H+ K+-6 TPage glaba 25 cubunit mBNA
9671	22323		0.56			EST HUMAN	J7837F Human feet heart, Lambda ZAP Express Home sapiens cDNA done J7837 6' similar to B-CELL RECEPTOR ASSOCIATED PROTEIN 78.00.00.00.00.00.00.00.00.00.00.00.00.00
9951			0.99	1.3E-01	393940	5	Rettus norvegious peptidy argining deliminase time IV (Data) mibrily
10030		35894	0.85	1.35-01	01 AW851599.1	EST HUMAN	MR2-CT0222-201099-001-601 CT0222 Home serilens cDNA
10291	25128	36151	1.1	1.3E-01		Π	Homo saplens chromosome 21 segment HS21C046
10423	23069	38290	0.64	1.3E-01		EST_HUMAN	AU121237 HEMBB1 Homo saplens cDNA clone HEMBB1002387 5/
10471	23117	36347	0.52	1.3E-01	_		2820637.3prime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2820637.3
10528	- 1		2.93	1.3E-01	2		MR4-BT0358-130700-010-h08 BT0358 Homo saplens oDNA
10//5	- 1	36701	1.58	1.3E-01		THUMAN	y32d09.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:150449 5'
11039	-	36980	1.33	1.3E-01	01 AF119117.1		Homo sapiens dopamine transporter (SLC8A3) gene, complete cds
11276	23879		3.28	1.3E-01	71745		Mus musculus cofilin 2, muscle (Cfi2), mRNA
11304	23963	37263	1.42	1.38-01			602087045F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4251348 5'
11304	23983	37264	1.42	1.3E-01			602087045F1 NIH_MGC_83 Hamo saplens cDNA clone IMAGE:4251346 5'
11589	24188	37504	4.26	1.3E-01			601158052F1 NIH_MGC_21 Homo septens cDNA clone IMAGE:3504804 5'
11723	24317	37640	1.94	1.35-01			601473369F1 NIH_MGC_68 Hamo sapiens aDNA done IMAGE:3876208 5'
11755	24346	37676	4.1	1.3E-01		. 1	602139760F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4300863 5'
12114	24607	31088	1.37	1.3E-01	1.3E-01 BE618346.1	EST_HUMAN (601462741F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866003 5'

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	Most Similer (Top) Hit Top Hit Acession Database BLAST E No. Source	1.3E-01 AJ242790.1 NT	4 2E 04 743004 4 NT	4 3E 04 AB026829 1		1.3E-01 AW001114.1 EST_HUMAN		1.2E-01 AI421744.1 EST HUMAN	1.2E-01 U66912.1 NT	1.2E-01 AF039442.1 NT	1.2E-01 AU149146.1 EST_HUMAN	1.2E-01 AU149146.1 EST_HUMAN	1.2E-01 AV735249.1 EST_HUMAN	1.2E-01 AA897474.1 EST_HUMAN		1.2E-01 Q14934	1.2E-01 AI285402.1 EST HUMAN	1.2E-01 X89211.1 NT	1.2E-01 AW 449368.1 EST_HUMAN	1.2E-01 BF248490.1 EST_HUMAN	1.2E-01 AL163213.2 NT	1.2E-01 AW998558.1 EST_HUMAN		1.2E-01 AI623388.1 EST_HUMAN	1.2E-01 U18018.1 NT		באוסער ופש	1.2E-01 M16364.1 NT	1.2E-01 X56882.1 NT	1.2E-01 AW370668.1 EST_HUMAN	1.2E-01 U67600.1 NT		1.2E-01 X56882.1 NT
	op Hit Acession No.	ŀ	Γ	B026829 1		W001114.1		1421744.1	J66912.1	(F039442.1	JU149146.1	(U149146.1	V735249.1	\A897474.1		114934	11285402.1	(89211.1				4W996556.1		AI623388.1	U18018.1		A1720470.1	M16364.1	X56882.1	AW370668.1	U67600.1	Z99118.1	X56882.1
-		_	-				_	ı		1.2E-01	1.2E-01	1.2E-01	1.2E-01/			1.2E-01	1.2至-01	1.2E-01	1.2E-01	1.2E-01	1.2E-0	1.2E-0											
	Expression Signal	4 43		10.1	?	2.26		8.42	1.05	4.33	3.22	3.22		1.23		1.26	2.81	2	1.03	1.75	1.2			1.12						1.59	1.12	0.8	0.82
	ORF SEQ ID NO:							25844			26778					27062	27082			27642	27748			28176	ł		28318	1 28350	28418		4	3	7 28921
	Exan SEQ ID NO:	SAROE	1		24915	95076	1	13199	<u> </u>		l		1	1	1	14373	L	L	Ĺ	L		1_	l	15438	L	ᆚ_	15659	15701	15770	15987	16014	16228	16267
	Probe SEQ ID NO:	1,004	147	122/4	12606	42R3R	202	374	416	534	1355	1355	1382	1408		1627	1646	1762	1913	2181	2284	2597		2731	7,80	150	2903	2935	3004	3224	3252	3472	3511

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		~		_	_			_				_			_				_											
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Wheat mRNA for a group 3 late embricogenesis ahi indani protein (IEA)	Bacillus subtilis complete genome (section 15 of 21): from 2705/31 to 3013540	P.clarkli mRNA; repeat region (ID 2MRT7)	P.clarkii mRNA; repeat region (ID 2MRT7)	Lesculentum mRNA for glyoxalase-I	Rana ridibunda pitultary adenylate cyclase-activating polypeptide variant 2 precursor, mRNA, complete cds, alternatively soliced	Inye3c04.s1 NCI_CGAP_GCB1 Homo septens cDNA clane IMAGE:12820503"	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively soliced	2008d02r1 Soeres parathymid frimor NhHPA Homo contract DNIA class Contract	Homo sepiens gene encoding plakophilin (expres 4.43)	M.domestica Borkh, Granny Smith adh mRNA for about downstone	601493518F1 NIH MGC 70 Home seniens CDNA clone MARCE 3805643 gr	MATING-TYPE P-SPECIFIC POLYPEPTIDE PI	ILO-CT0031-221099-113-e04 CT0031 Homo saniens cDNA	Mouse galactosyltransferese mRNA complete cds	nx85c01.s1 NCL CGAP GCB1 Home seniens cDNA close IMAGE:1280024.3	602023112F1 NCI_CGAP_Brn67 Homo sablens cDNA clone INAGE-1458988 R	JC virus agnoprotein, VP2, VP3, VP1, large T antiden and small antiden canaes complete and	PM3-BN0137-290300-002-f09 BN0137 Homo sapiens cDNA	wc99g03.X1 NCI_CGAP_Cc3 Homo sapiens cDNA clone IMAGE:2326804 3' similar to SW:GST2_HUMAN O99735 MICROSONAL CL) ITATHIONE C. TO ANGEED A CT.	NADH-UBIQUINONE OXIDORED ICTARE BOX SUBINIT/COMBIENT DOWN OF BOX	8t71b10.x1 Berstead colon HPLRB7 Home senties cDNA clare IMAGE 327732 21	xx49d07.x1 NCI_CGAP_Eso2 Homo sapiens cDNA clone IMAGE:2687697 3' similar to gb:M13452 LAMIN A (HUMAN);	Staphytococcus aureus plasmid pSK23 putative recombinase Sin (sin) cene, partial rds: and transcrinting	regulator QacR (qacR) and multidrug efflux protein QacB (qacB) genes, complete ods	N.crassa vacuolar ATPase 57-Kd subunit (vme-2) gene, complete ads	N.crassa vacuolar ATPase 57-Kd subunit (vma-2) gene, complete cds	Homo sapiens Xq pseudoautosomal region; segment 2/2	Haemophilus influenzae Rd section 29 of 163 of the complete genome
gle Exon Pro	Top Hit Database Source	TN	NT	N	FZ	Ę	Ā	EST HUMAN	Į Į	EST HUMAN	LN	LN LN	EST HUMAN	SWISSPROT	EST HUMAN	NT	EST HUMAN	_	FN	EST_HUMAN	EST HUMAN	Т	Т							LN L
Sin	Top Hit Acession No.	-01 X56882.1	-01 299118.1			-01 Z48183.1	01 AF221633.1	01 AA744369.1	01 AF223391.1				6.1	01 P10842	76.1	01 M26925.1	Γ	01 BF347985.1	1 AF295739.1	01 BE007072.1		Γ	1.2E-01 AI832681.1			2.1				1.2E-01 U32714.1
	Most Similar (Top) Hit BLAST E Value	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.28-01	1.2E-01	1.2E-01 Z48234.1	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2€-01 /	1.2E-01		1.2E-01	1.2E-01 J03956.1	1.2E-01 J	1.2E-01	1.2E-01
	Expression Signal	0.82	1.46	1.97	1.97	1.1	0.92	18.0	÷	2.59	2.3	0.68	1.83	1.36	2.35	1.64	0.57	1.14	0.69	1.4	4.36	0.67	0.73	. 10.29		3.34	0.99	0.99	1.09	1.49
	ORF SEQ ID NO:	28822			-	30041		30492	30649	30659	30749	30926	31853	31898	31960	32020	32097	32321	32658		33683	33731	34037				34184	34185		1
	Exen SEQ ID NO:	16267	16228	16900	16900	17406	17471	17979	18025	18034	18089	18217	18884	18930	18981	19043	19107	19315	19616	20488	20557	20601	20902	20989		21008	21047	21047	21191	21281
	Probe SEQ (D NO:	3511	3692	4160	4160	4672	4739	6170	6217	6227	5284	5418	6107	9183	929	6270	6337	8550	6700	7783	7862	7906	8208	8295		2 2	804	8364	8488	8589

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	M.musculus DNA fragment of Apolipoprotein B gene	S.cerevisiae HXT5 gene	AV710857 Cu Homo sapiens cDNA clone CuAAKEUS 5	601900763F1 NIH_MGC_19 Homo sapiens cDNA clone IMACE:4130103 0	Yeast MPT5 gene for suppressor protein, complete cds	601655578R1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3846283 3	601900763F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4130103 o	Homo sapiens dynein intermediate chain DNAI1 (DNAI1) gene, exon 1/	yf80c02.s1 Soares Infant brain 1NIB Homo saplens cDNA cione IMACE. 20000 3	Rabbit glycogen-associated protein phosphatase regulatory subunit (RG1) mKNA, complete cas	AV658033 GLC Homo saplens cDNA clone GLCFIB12 3'	Home seniens Xa pseudosutosomal region; segment 2/2	MACROPHAGE-STIMULATING PROTEIN RECEPTOR PRECURSOR (MSP RECEPTOR) (P185-RON)	(CDW136) (CD136 ANTIGEN)	Drosophija melanogaster strain Oregon R potential RNA-binding protein gene, complete cds, and syntamin	gene, partial cds	Home sapiens colon cancer antigen NT-CV43 IIINNA, parter cos	R.norvegicus NF68 gene for 68kDa neuroniament	QV4-BT0234-111199-031-910 BT0234 Homo sapiens cUNA	dn20g05.x1 NCI_CGAP_Lu5 Home sapiens cDNA clone IMACE:1898840 3	Xenopus laevis integrin alpha 3 subunit mRNA, partial cds	T	T	Т	Mus musculus protein (16kDa) similar to human SYK interacting protein (p16K), mRNA	Т	T		Г	Г	Т	Т	Т	Т	
Single Exon Probe	ORF SEQ Expression (Top) Hit Top Hit Acession (Top Hit Database ID NO: Signal BLASTE No. Source	0 62 1.2E-01 X15191.1 NT	1.2E-01 X77961.1 NT	4 50 4 2E-04 AV710857 1 EST HUMAN	1 2F-01 BF314481.1 EST HUMAN	NT N N N N N N N N N N N N N N N N N N	1.2E-01 D20184:1	1 2E-01 BE314481 1 EST HUMAN	1.30 1.4E-01 D1.31 NT	NAMUM FIST HUMAN	STOCK MEETON 4	1.ZE-01 MOSTOS:	1.2E-01/AV608033,1		SWISSPROT	2,010	3 16 1.2E-01 AF188892.1 NT	1.2E-01 AF039442.1 NT	1 2F-01 X53981 1 NT	NEW PERSONAL FEST HUMAN	1.44 1.2E-01 BE001410:1	1.2E-01 Atabados 1	1.ZE-UTICIOIOF.I	7.95 1.2E-01 (080455)	1.ZE-01 AEU04428.1	1.81 1.2E-01 289110.1	1.38 1.4-01	T	4 65 1 1E-01 AA569006.1 EST_HUMAN	33.	1 1 04 N 1 44 8 8 0 0	1.3 1.1E-UTIALIGISOU.2	4.62 1.1E-01 AW9/2130.1 EST_CIONARY	3.01 1.1E-01 D64004.1	AU140363.1	1.1E-01
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Table 4

	Top Hit Descriptor		11), mRNA	saplens cDNA	HSC1RF022 normalized Infant brain cDNA Homo saplens cDNA clone c-1rf02 3'	Mus musculus calcium channel, voltage-dependent, T type, alpha 1G subunit (Cacna1g), mRNA	OU 3085/9F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3627066 5'			VEXIN 50) (CAP-50)	Cugalius gene encoding non-histone chromosomal protein HMG-14b, excus 4 and 5	sapiens cDNA	saplens cDNA	mRNA, complete cds	Tapa-1=Integral membrane protein TAPA-1 (mice, B cell lymphoma line 38C13. Genomic 1973 of segments	ווים ווים ווים ווים ווים ווים ווים ווים		Mus musculus major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyt transferase-alpha, palmitoy-protein thioesterase 2 (PPT2)		INTERPORT OF THE PROTEIN 3 (SCP-3 PROTEIN) INTRA03.51 NC CGAP EW1 Homo saplens cDNA clone IMAGE-1788440 cloneling to	ent;	gene, exon 6	ditions of nitrogen deprivation				yens cDNA	piens cDNA	182	24 323 45 Tullitan salivary gland cell line HSG Homo sapiens cDNA clone RL43	ACETYL-COENTYME A SYNTHETAGE A CONTRIBUTION OF THE CONTRIBUTION OF	HOME SOURCE THE LASE (ACEIAIE—COA LIGASE) (ACYL-ACTIVATING ENZYME)
Single Exon Probes Expressed in Brain	Top Hit Database Source	77 (1	Т	Т	ES I HUMAN HSC1KF022 nomalized infa	NI Mus musculus calcium chant	7	C.remardti nuclear gene on linkage group XIX	CWISCODOT ANNIESTS SENETOR VANSOSASE	Т	EST LINAM MADS STORES COLORS	Т	Т	EST HIMAN III & IMMOTO 2000 ass 11,0000 in the complete cds	T	'dı / j	A.immersus gene for transposase	Mus musculus major histocom Notch4, PBX2, RAGE, Iysopho	SWISSPBOT SYNADTONEM COMPLEX COMPLEX		HUMAN	6 Homo sapiens diacylglycerol kinase 3 (DAGK3) gene, exon 6	Douyus cinerea strain 14 cDNA library under conditions of nitrogen deprivation	S. pormbe sues gene encoding protein kinase	rowdencia rettgen penicilin G amidase gene	FST HIMAN DAYS ETONS 420000 500 500 500 500 500 500 500 500 50	Т	Т	FST HIMAN AF035748 Limes Ciliamosome 21 segment HS21C082	Т	Т	Т
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	Expression Signal	-	1 17	0.82	1.76	2.11	121	0.77	0.74	1.47	1.28	1.28	8.78	0.77	96.0	12	!	0.85	-	;	1 8	0.58	1.81	5.31	1.67	1.75	9.4	0.67	1.25	0.72	3.71	3.03
	ORF SEQ ID NO:			28442	L	28822				29089	29456	29457	-	29624	29956	30134			30412		31381	31399	31459	31500	31676	31696	31719	32088	32098	32149	32278	_
	Exon SEQ ID NO:	15530	15286	15796	16096	L		16325	16441	L	16833	16833	16967	16995	17329	17612		16839	17796	18381	18448	18480	18537	18572	18718	18737	18757	19100	19108	19150	19277	19367
	Probe SEQ ID NO:	2540	2572	3030	3336	3415	3444	3570	3688	3696	4090	4080 080	4228	4264	4694	4780		4957	2077	5584	5863	2882	5745	5781	5936	5955	5975	6330	6338	6381	6512	6604

Page 113 of 536 Table 4 Single Exon Probes Expressed in Brain

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	Expression Signal	275	3.91	1.39	1.31	1.58	3.21	3.21	2.81	1.65	1.36	3.92	1.97	3.15	2.74	2.63	1.95	1.1	1.19	1.03	96.0	96.0	2.53	0.95	1.19	2.35	9.73	1.21	14.15	0.99	0.7
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Page 115 of 536 Table 4 Single Exon Probes Expressed in Brain

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Table 4
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Single Exon Probes Expressed in Brain	Top Hit Descriptor		Drosophila malanorasia- ftz none	601065554F1 NIH MGC 10 Home parities - PNA -	Social Services of the Color of	Sacutaromyces cerevisiae suppressor of ABF1 (SAB2) gene, complete cds Bacillus haloditrane generals DNA	Drosophila melanogaster cAMP-dependent protein kinase type II requisitory enhinit (Av. Din Dana	complete cds	601070219F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3456385 5	601070218F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3456385 Fi	Homo saplens neurexin III-alpha gene, partial cds	Zuesco3 x5 Sceres overy tumor NbHOT Homo sepiens cDNA clone IMAGE:740932 31	/d//c12.x1 NCI_CGAP_Lu24 Home septens cDNA clone IMAGE:3278998 3'	Aspergillus terreus BSD mRNA for blasticidin S deaminase, complete cds	repetitive element contains element MIR MIR repositive element contains Alu	xd43c09.x1 NCI_CGAP_Ov23 Homo sapiens cDNA clone IMAGE:2896628 3' similar to contact and	repetitive element, contains element MIR MIR repetitive element :	Mus musculus phospholipid transfer protein (Plip), mRNA	U.sativa KAmy3C gene for alpha-amyfase	Homo sapiens I factor (complement) (IF) mRNA	Usucus carota leucoanthocyanidin dicxygenase 2 (LDOX) mRNA, LDOX-2 allele, complete cds	Leptosphaeria macuians beta-tubulin mRNA, complete cds	Leptosphaeria maculans beta-tubulin mRNA, complete cds	ruman ne i P deta mKNA for protein tyrosine phosphatase delta	Fruman laminin B1 chain gene, exon 26	PULYSOLYSST I NIH_MGC_66 Home sapiens cDNA clone IMAGE:3864287 5	Nattus norvegicus microtubule-associated protein tau (Mapt), mRNA	Aloe arborescens mRNA for NADP-mailc enzyme, complete cds	Homo sapiens fibroblast growth factor recentor 3 (cohond-color)	QV1-HT0518-070300-005-004 HT052-1-0000000000000000000000000000000000	CELL SUBSACE A 22 ANITORIA IN TO NOT BE SERVED SOLVA	Caulobacter crescentus thymydilate kinase (trik) and DNA polymerase III delta prime subunit (dnaC) canae	Complete Cds
igle Exon Pro	Top Hit Database Source		뉟	EST HUMAN	N 12	LZ.		I N	EST HUMAN	ES HUMAN	Z	EST HOMAN	ES TOMAN	Z	EST_HUMAN		EST HUMAN			N.	I L	-IN	±N.		T LILINAAA!	NEMOL				EST HUMAN	T		
ਲੋਂ -	Top Hit Acess		-01 X00854.1	:-01 BE537719.1	1.0E-01 U66834.1	1.0E-01 AP001507.1	9 9E-02 A E274008 4	0 0E 02 DEE18FE11	9.9L-72 DE340094.1	0 0E 02 A E000040 J	9.9E-02 AFUBBB10.1				-02 AW103088.1		02 AV 103088.1		4504570	02 AF484274 4	T	T	T		-	9754	9 7E-02 AROOFBOB 4		4503710 NT		9.7E-02 Q99795		
	Most Similar (Top) Hit BLAST E Value	10,	1.0E-0.	1.0E-0.	1.0E-01	1.0E-01	9.9E-02	20.00	0.9E-02	0 000	9.9E-02	9 9F-02	9 9F-02		9.9E-02	0.05.00	0.0E.02	9 8F-02	9.8E-02	9.8F-02	9.8E-02	9.8E-02	9.8E-02	9.8E-02	9.8E-02	9.8E-02	9.7F-02/4	77	9.7E-02	9.7E-02 E	9.7E-02/C	9.7E-02 A	
	Expression Signal	200	77.7	2.74	5.03	9.8	1.27	1 53	153	133	0.75	0.93	9.17		0.93	0.93	88	1.43	60	3.64	6.24	6.24	0.77	1.18	2.05	1.78	1.31		1.49	2.08	3.48	0.94	
	ORF SEQ ID NO:						28224			28671	29324	30003	30547		33634	33635	35065			28550	29564	29585			36436		26752		,0220	5//24		30695	
	SEQ ID NO:	24738	1	⊥	_[25002	15486	15495	L	L	l i		17951	- 1	20510	20510	21887	13333	15865	15906	16939	16939	20091	21884	23204	24570	14077	9257	14004	502	16/14	18067	
	Probe SEQ ID NO:	12317	1	47021	/071	12733	2781	2790	2790	3260	3933	4632	6875	407	(813	7815	9156	220	3100	3142	4198	4198	7381	9153	11437	12052	1328	1580	2257	1000	CDA	5261	
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Page 117 of 536 Table 4 Single Exon Probes Expressed in Brain

					Juin	le Exon Piou	Single Exon Probes Expressed in Drain
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
							Caulobacter crescentus thymydilate kinase (tmk) and DNA poymerase III deta piirite succiin (missy server)
5261	18067	30696	0.94	9.7E-02	AF099189.1	- 1.	complete cds
5924	_	31662		9.7E-02		EST HUMAN	ES 380348 MACE Lescapane (section 16 of 21); from 2997771 to 3213410
7198	1	32958	3.24	9.7E-02		Z	Decillus suuria von promo german Officerov Foithellum Homo saplens cDNA clone IMAGE:254788 3'
7882	L	L				EST HUMAN	W41003.51 Wezirrain Oractory Epithelium Homo sapiens cDNA clone IMAGE:254788 3'
7882	L		1.28	9.7E-02	N22798.1	EST HUMAN	W44 1003; SI WELTHRING CLEAP ON AR Home sepiens cDNA clone IMAGE: 2549747 3' similar to gb: X52851_me1
8748		34587	1.49	9.7E-02	Al953984.1	EST_HUMAN	PEPTIDYL-PROLYL CAST AND SCHOOL (HUMAN);
11152	L		2.84	9.7E-02	U58337.1	Z	Mus musculus ligatin (1940) ilinviv, per use accession of the light of
2000	L	27470		9.6E-02	AI080721.1	EST HUMAN	oz47d11x1 Soares NnHwiPu_SI Homo septents CDNA clone IMAGE:16784853'
2000			1.11		Al080721.1	EST_HUMAN	loz47d11x1 Soares unmiwing of Holling saperito Commentation of the Comment of Holling Saperito Comment of the C
4311	L	L	5.8	9.6E-02	Z32686.2		Proteus mirabilis timutia operuri, surair i inversione con A
4040	L	L	0.99	L	AW966230.1		ES 1378303 MAGE resequences, which i toling deposit of the second of the
6014	L	l	3.13		BE910039.1	EST HUMAN	601498086FT NIT MOC / OTTAIN SERVING CONE PLACE1005740 5
8274	L	8	9.0		AU137084.1	EST_HUMAN	AUTS/1064 FLACE Indites agricus and Artificial CACAAHO2 5
	1	35300	1.31			EST HUMAN	AV68/686 GNC Home saprens CDNA clone IMAGE:3919363 5
776	1	23	1.12	9.6E-0.	2 BE894895.1	EST HOMAN	801434060F11Nii MCC - Francisco Communication Communicatio
6586	1	35790		9.6E-02	2 AJ243211.1	Į.	Hame sepiens DMRT1 candidate tumour suppressor gene, exons 1 to 55
88	ı		,		2 AJ243211.1	N. P. S. S. S. S. S. S. S. S. S. S. S. S. S.	Anna REFERENTIA MGC 83 Homo septens cDNA clone IMAGE:4250969 5'
1002		35884	4 0.5		9.6E-02 BF677270.1	ES HOMAN	A HATHINITY mail is transposon Tem3 bseudogene for transposase (in S-5 copy)
1005	1				2 AB013985.1	Z	Anuminimum majus transposon Tam3 pseudogene for transposase (in S-5 copy)
10051	L	35916			9.6E-02 AB013985.1	TO00001110	COMBI EMENT DECAYACCELERATING FACTOR PRECURSOR (CD55)
10158	38 22806				9.6E-02 P08174	DE LA SERVICIO	MAnchacterium tuberculosis H37Ry complete genome; segment 102/162
10643	L	34 36572			9.6E-02 Z79702.1	NOT INVOID	without 3 Sogres Infant brain 1NIB Homo sapiens cDNA clone IMAGE: 48653 3
12652	52 24954		e.		9.6E-02 H14589.1	TOT LINAN	CM2-BN0023-050200-087-f12 BN0023 Homo sapiens cDNA
4081	16825	25 29452			9.5E-02 AW992395.1	ESI HOWAN	TRANSKETOLASE 2 (TK 2) (TRANSKETOLASE RELATED PROTEIN)
5579	79 18376	76 31289	9 0.85		9.5E-02 P51854	DATE NOT	organo e1 Stratagene fetal retine 937202 Homo saplens cDNA clone IMAGE:867736 3
9869	l_	81 32729			9.5E-02 AA780728.1	EST HUMAIN	Trimograminate flexibilities DNA for phospholipase A2 inhibitor, complete cds
7202	L	88 32963			9.5E-02 AB003473.1	Z	A radiancia theiliana DNA chromosome 4, contig fregment No. 38
7467	67 20141	41 33234			9.5E-02 AL161538.2	2	TRANSPETCH ASE 2 (TRANSKETOLASE RELATED PROTEIN)
7597	L	76 31289			9.5E-02 P51854	SWISSPROI	F07453642F1 NIH MGC 66 Homo sepiens cDNA clone IMAGE:3857243 5
7,88	80 20475	75 33600			9.5E-02 BF035861.1		RATATSA27F1 NIH MGC 66 Homo sapiens cDNA clone IMAGE:3857243 5
7780	L			١	9.5E-02 BF035861.1	EST HUMAN	601453642F1 NIH MGC 66 Hamo saplens cDNA clane IMAGE:3857243 5
10578	1	73 36509	2.36		12 BF035861.1	EST_HUMAN	101-121-121-121-121-121-121-121-121-121-
2				١			

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Single Exon Probes Expressed in Brain	Similar Top Hit Acession Database Top Hit Acession STE No. Source	02 BF035861.1 FST HIMAN	02 BF671063.1 EST HIMAN	02 U55944.1	02 U55944.1 NT	02/233059.1	02 AF097363.1	7.46883 4 NIT	02 L78833.1 NT	22 U31815 1	2 AF198036 1 NT	22 4809280 NT)2 6912525 NT	12 BF575511.1 FST HIMAN	2 BE391943.1 EST HUMAN	2 BE391943.1 EST HUMAN		2 AP001507 1 INT	2 AW568007.1 FST HIMAN	2 BE962631.2 EST HIMAN	015034	215034 SWISSPROT	EST_HUMAN	FN	9.3E-02/AW468850.1 EST_HUMAN hd28h12.X1 Soares NFI T GBC ST Homo remains a Final statement of the statement	ļ	galactosyl transfer as (beta1.3-galactosyl transfer as (beta1.	2 2	U60315.1	R54156.1 FST HIMAN	Q28631 SWISSPROT	AA534354.1
Sin		Ī				Ī	Γ				-	08280	6912525IA		Ī	l				T		Γ					T	T				T
	Most Similar (Top) Hit BLAST E Value	9.5E	9.4	9.4	9.4	9.4E-02	9.4E-02	9.4F-02	9.4E-02	9.4E-02	9.4E-02	9.3E-02	9.3E-02	9.3E-02	9.3E-02 L	9.3E-02		9.3E-02	9.3E-02	9.3E-02	9.3E-02	9.3E-02 C	9.3E-02 A	9.3E-02 A	9.3E-02 A	0.35.02	9 2F-0211	9.2F-02[[]	9.2E-02 U	9.2E-02 R	9.2E-02 Q	9.2E-02 A
	Expression Signal	2.36	2.82	0.99	66.0	4.43	0.63	2.48	2.44	1.78	1.92	1.97	6.32	1.85	3.51	3.51	2.04	0.67	0.62	2.15	3.67	3.67	3.5	2.51	8.43	6	8.37	8.37	8.37	1.68	3.28	0.85
	ORF SEQ ID NO:	36510			L				33258					28665	29502	29503			33972	35449	35953	35954	+	1		 -	25668	25669	25670	-	28587	28709
	Exon SEQ ID NO:						18999	21180	20166	25255	24965	15754	15792	16013	16874	16874	17419	18373	20840	22263	22739	22739	2000	19167	25209	25254	13034	13034	13034	14952	15938	16061
	Probe SEQ ID NO:	10578	1825	1857	1857	3860	6225	8488	10851	11941	12671	2988	3026	3251	4132	4132	4685	9226	8148	9610	1009 P	600	2120		2020	12752	222	222	222	2224	3175	3289

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	Top Hit Descriptor	Nice miscripts are Tidel antiden receptor alpha (Ptora), mRNA	Mus muscaus pro commercial attention associated transcript, promoter region	Annotation of the MGC 17 Homo sapiens cDNA clone IMAGE:2960176 5	Statement of Statement (#837225) Homo sapiens cDNA clone iMAGE:69808 5 similar to similar to	JASSOCIATI STREET IN THE STREET (HUMAN) BEXEGOOG GUANINE NUCLEOTIDE-BINDING PROTEIN G(S), ALPHA SUBUNIT (HUMAN)	H.vulgare xylose isomerase gene	Mesocricetus auratus oviductin precursor (OVI) gene, compress cos	Podospora anserina mitochondrion, complete genome	O. cuniculus K12 keratin gene	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 54	A TAKE SEC. THE SEC.	Homo sapiens MSH55 gene, partial ods; and CLIC1, DDAH, G6b, G6c, G5b, G6d, G6e, G9t, BA 15, G5b, G6c, G6c, G6c, G7t, BA 15, G5b, G6c, G6c, G7t, BA 15, G7t, G7t, L7T, LTB, TNF, and LTA genes, complete cds	24-05 A Schmelder fettel brain 00004 Homo sapiens cDNA done IMAGE:2781968 5	41/4803.3 Scrimence Company Acceptance Activities the management of the Complete cds	Mus musculus digitarporani zon min	FB10F10 September Stratagene Homo saplens cDNA clone FB19F10 3'end	rojet vivi cara prami tana estas gratila≃sea urchins, embryos, Genomic, 5275 nt]	1go 10=Cyl acutil 11throades 8:	Aufahanana Shahanana musale 937209 Homo sapiens cDNA clone IMAGE:611783 3' similar to	SW:TRT3 HUMAN P48378 TROPONIN T, FAST SKELETAL MUSCLE, ISOFORM BETA:	Rettus norvegicus cell cycle protein poocuru gaile, compress con	Homo sapiens partial MUC3B gene for MUC3B finding states 1-11	FOLATE RECEPTOR ALPHA PRECURSOR (FR-ALPHA) (FOLATE RECEPTOR ALPHA PRECURSOR (FR-ALPHA) (FRED) (OVARIAN TUMOR-ASSOCIATED	RECEPTOR, ADULT) (ADULT) h/39g10.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3175842 3' similar to contains Alu	repetitive element	HIV-1 pscuss-no main OSA envelope strong envision (env.) gene, partial cds	PIN-1 pocusodo nom con constanta protein SP65 (cotE) gene, complete cds	Distriction discondent sport of These 3 gene	Hastington action of the second of the secon			
The state of the s	Top Hit Database Source			7	NUMBER TO	EST_HUMAN 9									HUMAN				EST HUMAN		LZ.	EST_HUMAN				TOddsolwo	Т	T_HUMAN	NT	LN.	LZ	N	
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Probe NO: NO: NO: NO: 1444 4214 4214 4214 11062	Exon SEQ ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:	ORF SEQ ID NO: 34719 37704 37709 377	Signal Signal Signal Signal Signal 3.55 0.99 0.99 0.99 0.97 0.57 0.57 0.57 0.57 0.57 0.57 0.57 0.5	Most S. Cap B. B. B. B. B. B. B. B. B. B. B. B. B.	Top Hit Acess No. No. No. A50 A750.1 A416720.1 BE264455.1 BE264455.1 A271881.1 A1167281.1 A1167281.1 A1167281.1 A1167281.1 A1271885.2 A721885.1 A721885.1 A721885.1 A721885.1 A721887.1 A72187.1 A721	Top Hit Source Source Source Source Source THUMAN THUMAN VISSPROT THUMAN VISSPROT THUMAN ST HUMAN ST H	Top Hit Descriptor TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT (TAFIL-135) (TAFII-130) TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT (TAFII-135) (TAFII-130) TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT (TAFII-135) (TAFII-130) Homo sapiens blood box gene 6 (aninfalls, keratitis) (PAX6), isoform b, mRNA Homo sapiens blood box gene 6 (aninfalls, keratitis) (PAX6), isoform b, mRNA Sheep mRNA for angiotensinogen, complete cds TRISBAGE 51 Strategere colon (#837204) Homo sapiens cDNA clone IMAGE:5566289 3' TRISBAGE 51 Strategere colon (#837204) Homo sapiens cDNA clone IMAGE:555648 6' TRISBAGE 51 Strategere colon (#837204) Homo sapiens cDNA clone IMAGE:355648 6' TOTOKINE INDUCIBLE SH2-CONTAINING PROTEIN 3 (PROTEIN EF-10) S.cerevisiae chromosome XIV reading frame ORF YNIL285W OCTOKINE INDUCIBLE SH2-CONTAINING PROTEIN 3 (PROTEIN EF-10) S.cerevisiae chromosome XIV reading frame ORF YNIL285W OCTOKINE INDUCIBLE SH2-CONTAINING PROTEIN 3 (PROTEIN EF-10) S.cerevisiae chromosome XIV reading frame ORF YNIL285W OCTOKINE INDUCIBLE SH2-CONTAINING PROTEIN 3 (PMCA3) gene, partial cds genes, complete cds; and plasma membrane calcium ATPase Isoform 3 (PMCA3) gene, partial cds Mus musculus and plasma membrane calcium ATPase Isoform 3 (PMCA3) gene, partial cds Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potassium channel protein, exons 10-14 Mus musculus partial Korroff gene for potass
10610	-1.1	3 37237			8.7E-02 AJ007763.1	N F	Gluconobacter oxydans tRNA-lle and tRNA-Ala genes Human DNA for immunoglobulin alpha heavy chain from a case of alpha heavy chain disease
12145					8.7E-02 X17146.1 8.7E-02 6679057 NT	TN 72	Mus musculus nidogen 2 (Nid2), mRNA Home eaplens Xq pseudoautosomal region; segment 2/2
1230 2240 3183	1230 13979 2240 14968 3183 15946	26649 27706 6 28596	9 7.02 6 1.82 6 4.57	8.6E	8.6E-02 AJ271/36.1 8.6E-02 BE408667.1 8.6E-02 L05468.1	EST_HUMAN NT	601304016F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3638643 5 Trichomonas vaginalis beta-tubulin (btub1) gene, complete cds

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	Top Hit Descriptor		e, complete cds	one IMAGE:4310259 3'		switch region (part a)	switch region (part a)	RSOR (IRR) (IR-RELATED RECEPTOR)	CAP) mRNA	CAP) mRNA	06), mRNA	nolog PrtC (prtC) gene, complete cds	nd unknown gene	one IMAGE:2972846 3'	omplete cds	Lacerta media cytochrome c oxidase subunit 1 gene, partial ods; mitochondrial gene for mitochondrial product		and a second mount of the second seco	ne IMAGE:4139216 5'	ne IMAGE:4139216 5'	Bacillus stearothermophilus BsrFI methylase (FIM) and BsrFI restriction endonuclease (FIR) genes, complete		oq83b07.s1 NCI_CGAP_Kid6 Homo sapiens cDNA clone IMAGE:1592917.3' similar to gb:K01144 HLA	A CHAIN PRECURSOR (HUMAN);		tiel cds		cDNA	cDNA		SZORF5), mRNA	ods
Single Exon Probes Expressed in Brain	± 83 op		Т	┰	T	Mouse germine 1gM chain gene, D region; D-452, mu switch region (part a)	7	Т	Home saperils oniz-related CBP activator protein (SRCAP) mRNA	Honey September Sontz-related CBP activator protein (SRCAP) mRNA	Pirtractilism 3	Universational discolateum proteasome subunit C2 homolog PrtC (prtC) gene, complete cds	T	Т	rames its vegicus of A-1 like protein p1294 mKNA, complete cds	Lacerta media cytochrome c oxidase subunit 1	I acerta media cipra a comprehensi l	Ť	Ť	Т	Badillus stearothermophilus BsrFI methylase (FIM) and BsrFI restri	Helicobacter pylori 26695 saction 130 of 134 of the complete	Г	Т	Т	Mis missilis phospholipase C-like protein mRNA, partial cds	Т	Т	Т	Homo sapiens chromosomo 22	Homo septens hendranges provided and traine 5 (C220RF5), mRNA	Complete of the control of the contr
gle Exon	Top Hit Database Source	1		ES! HUMAN			SIA/ISCODOT		1		Į.	Į.	TOT LINA	TN TOWN		L	Ž	FST HIMAN	EST HIMAN	LN	E	N	NAMIN TSE	TORIGORIANS	IN		ST HIMAN	EST HIMAN				
Sin	Top Hit Acession No.	02) AE1 533 50 4	l					5730066			U60168	23		Γ		6.0E-02/AF-206551.1	8.6E-02 AF206551.1	Γ		8.6E-02 AE001073.1		8.5E-02 AE000652.1	8.5E-02 AA985491.1		35.1	6754779	BE833054 1	Γ	Γ	1418108	8.5E-02 AF155510.1 NT	Ī
	Most Similar (Top) Hit BLAST E Value	A SF.	9 9	8.6E-0	S GE	8.6E-0	8.6F-0	8.6E-02	8.6E-0	8.6E-02	8.6E-02	8.6E-02		8.6E-02	1 20 0	8.05-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.6E-02	8.5E-02	8.5E-02	8.5E-02 P08089	8.5E-02	8.5E-02	8.5E-02	8.5E-02	8.5E-02	8.5E-02	8.5E-02	0 57 00
	Expression Signal	3.77	98	4.75	1.58	1.56	1.34	1.25	1.25	0.62	0.81	1.76	0.58	0.81	0	2	1.8	4.64	4.64	5.97	2.11	3.3	0.75	1.29	6.95	1.65	2.81	2.81	0.54	0.87	8.87	67 7
	ORF SEQ ID NO:			31746	32033	32034	33248	33651	33652	33788		35482		35918	36702	3	36793	37163	37164	36414	37481	27853	31292		31658	34340	35591	35592	36119	36243		37070
	Exon SEQ ID NO:	16388	17852	L	L	Ĺ.	20153	L		20664	20726	22289	22326	22701	23545		23545	23877	23877	23184	24167	15116	18380	18417	18706	21196	22387	22387	22909	23028	23775	23794
	Probe SEQ ID NO:	3635	5134	6003	6281	6281	7481	7831	7831	7969	8031	9637	9673	10053	10865		10865	11214	11214	11417	11568	2396	5583	2621	5921	8504	9736	9736	10261	10382	11105	11125

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Top Hit Descriptor EST72736 Ovary II Home saplens cDNA 5' end EST72736 Ovary II Home saplens cDNA 5' end ad441.1.4 Soares_fetal_beart_NIHHH9W Home saplens cDNA clone IMAGE:2350221 3' similar to contains element i. ad441.1.4 Soares_lost_in beart_NIHH9W Home saplens cDNA clone IMAGE:2350221 3' similar to contains element ii. Cavia porcellus gi/coproficial alpha-subunit ImRNA, complete cds element MSR1 repatitive element i: Cavia porcellus gi/coproficial alpha-subunit ImRNA, complete cds Home saplens mRNA for FLUDIODE protein, Paul and idea in the capiens cDNA clone IMAGE:3534393 5' Home saplens mRNA for FLUDIODE protein, Paul and idea in the capiens cDNA clone IMAGE:3534393 5' Home saplens mRNA for FLUDIODE protein, Paul and idea in the capiens cDNA clone IMAGE:145895 5' Home saplens mRNA for FLUDIODE protein capiens cDNA clone IMAGE:215210 3' MSSN12.11 Soares placenta NICATORIONICO, complete gardene Ixodes hexagonus milcohoridino, complete condition subjetes cDNA clone IMAGE:2152103' Wordert Ixone Sazlenes INOI_CGAP_Kdd Home saplens cDNA clone IMAGE:2152103' MGSTATA Sazles INOI_CGAP_Kdd Home saplens cDNA clone IMAGE:215322 015332 GAMMA SuBJUNIT OF SODJUM POTASSUM Apr PASE LIKE: Gallus gallus mRNA for for OBGAP more pagenes cDNA clone IMAGE:39299935 5' Gallus gallus mRNA for for OBGAP more pagenes cDNA clone	THUMAN TH	Top Hit Aces No. No. No. No. No. No. No. No. No. No.		Signal Signal 3.8 3.73 3.74 4.97 7.17 1.07 1.08 3.09 2.98 4.1.72 1.09 1.09 1.09 1.09 1.09 1.09 1.09 1.09	ORF S S S S S S S S S S S S S S S S S S S		Probe NO: 12700 2672 3801 4321 4321 4321 4321 6530 7028 7028 10260 12070 2006 3507 2006 3507 12070 12158 1157 1157 1157 1157 1157 1157 1157 1
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Abdone the line a DNA chromosome 4, contig fragment No. 91	Т	2 AW583503.1					
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and A Baretoned colon HPLRB7 Homo sapiens cDNA clone IMAGE:2335842 3' similar to 1 K: U86314		AF218890.1	8.4E-02			L	8741
no semiens attractin precursor (ATRN) gene, exon 2		BEOSOV4.1	8.4E-UZ			L	7928
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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Law emiser	From Septems of formosome 21 segment HS21C006	LEUCOCYTE ANTIGEN CO97 PRECURSOR	LEUCOCYTE AN TIGEN COBY PRECURSOR	Mile milecture of the contraction of the contractio	This interculus Jin valisporte (Jn -3) gene, complete cds	FOLD ASSESSED IN THE MACHINE M	Boctomore Mind MicC_/Z Hamo septems cDNA clone IMAGE:3924523 6	AV743344 OB U.S. Complete dissue growth factor precursor (CTGF) gene, complete cds	ROS BTOOK ASSESS OF THE PROPERTY OF THE PROPER	Rock north culture of the Prince Sapiens cDNA	Devenion of the patron with VIDS KNA-2	Chlam: Joship and 1.1. A Page Splens cDNA clone IMAGE: 3366696 6	Mis misculin addoction of the complete genome	alternatively shillson and complete cds. (Egfr) gene, exons 5 through 28, and complete cds.	William of July Company of 2000 of the Company of t	A A A MARKET LANGUAGE OF THE COMPINED GOVERNOON OF THE COMPINED GOVERN	Litter result romo sapiens culva clone A1484	indicated a first conformation of the second	Moodloc.X1 NCL_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2338603 31	nomo sapiens hypothetical protein FLJ10060 (FLJ10060), mRNA	Homo saplens hypothetical protein FLJ10060 (FLJ10060), mRNA	Home sapiens extracellular glycoprotein facrifin precursor, gene, complete cds	FST368723 MADE securities 11 Segment HS21C002	Molluscum contactorium virus eurobiano (2000)	Human dene for dibudraling and designed in complete genome	Himen gene for diriyu iliyodiila succiniyilahishrase, complete cds (exon 1-15)	The state of the s	Synechowstis on Picked committee of the september of the	Symptocytics of the second confidence generated 1/12/, 213/258-2267269	777269-701	Dichaselling discalaring and a september CDNA clone IMAGE:4076619 5'	Thermonlasma and orbitime control of the phosphodiesterase gene, complete cds	EST378191 MAGE resequences. MAGI Homo serviens clinia	Ultra circular
gle Exon Pro	Top Hit Database Source	ŀZ	CWICEDEAT	SWISSPACI	SWISSEDER	NT NT	LZ	EST HUMAN	IN LA	EST HIMAN	19		TOT HIMANI	TOWN TO LA		Ż	LZ	FST HIMAN	LA LA	EQT UPLAN	NCWOL - S		- E	L	ST HUMAN	N.		LZ.	HIMAN	Т		HIMAN			T_HUMAN	
Sin	Top Hit Acesslon No.	8.2E-02 At 163206 2	P4896n	8.2E-02 P48960	P48960				ı	8.2E-02 AV743341.1	T	Τ	-				8.1E-02 AE004006.1		2	T	26074	44 4000744	02/AY005150 1	T					-			-		-		
	Most Similar (Top) Hit BLAST E Vatue	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2至-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02		8.2E-02	8.1E-02	8.1E-02 T115321	8.1E-02	8 1E-02	8.1E-02	8 4E 02	8.1E-02/	8.1E-02	8.0E-02/	8.0E-02	8.0E-02 D26535.1	8.0E-02	8.0E-02.B	8.0E-02	8.0E-02 D	2	12	8.0E-02.A	8.0E-02 A	
	Expression Signal	1.07	4.97	4.97	4.97	2.44	2.39	1.49	3.09	29.0	2.95	5.36	2.24	4.03		3.65	0.79	1.19	0.66	1.26	0.61	0.84	18.	2.08	5.03	0.79	9.85	9.85	3.27	1.09	1.09	4.2	. 0.98	1.45	1.01	
	ORF SEQ ID NO:	29371			29619				32707		34509	35332	35512	31102			31378	32040	-	-	34067	34068	-	37396	25447	26344	27134	27135	27343	27835	27836		26486	28315	29182	
	SEQ ID	16737	16992	16992	16992	17743	ı	ı	- 1			- 1	22315	24646		25138	18483	19069	19786	20154	20932	20832	22463	24083	15534	13682	15576	15576	14633	15096	15096	15191	13827	15687	16549	
	Probe SEQ ID NO:	3989	4261	4251	4251	6022	5070	5252	6925	7632	8670	9499	9663	12164		12664	2008	988	7097	7482	8238	8238	9812	11482	9	916	1694	1694	188	23.74	2374	2473	2823	2304	3797	
														_									_			_		_	_	_	_	_		_	—	

Page 125 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	AND WALL OF AND Gas Home sapiens CDNA clone IMAGE:2132114 3'	IBOEXI NOTOGO	M.musculus gene for genericase by the Demonstration of the MAGE:2721547 3'	H-Bit-fid-f-10-0-U. st NCI_CGAP_Subs name squeris constitution	Homo sapiens ABCA1 (ABCA1) gene, complete cds	Homo sapiens ABCA1 (ABCA1) gene, complete cds	Botrytis cherea strain T4 cDNA library under conditions of nitrogen deprivation	H.sapiens AGT gene, intron 4	H saplens AGT gene, Infron 4	Homo septens chromosome 21 segment HS21C009	Homo sapiens SCG10 like-protein, helicase-like protein NHL, M68, and ADP-ribosylation factor related	protein 1 (ARFRP1) genes, complete cds	Drosophila orena hunchback region	Homo saplens cAMP responsive element birding bruentance (10 to 10	600943191F1 NIH MGC 15 Hamo sapiens culva cione invasti consociale de la c	ar Berstead colon HPLRB7 Homo septens cDNA clone IMAGE: 2173646 3' similar to go: 2000 o	60S RIBOSOMAL PROTEIN L38 (HUMAN);	Plasmodium falciparum strain Dd2 heat shock protein 86 (HSP86), O1 (o1), O3 (o3), U2 (o2), CG3 (cg3),	CG4 (cg4), CG3 (cg3), putative chloroquine resistance transporter (crt), CG9 (cg9), CG1 (cg1), CG5 (cgo,	CG2 (cg2), and CG7 (cg7) genes, complete cus	Mus musculus colony sumulating factor 1 receptor (Ceffr), mRNA	MUS MUSCAULS CAGAD SINICIPATION PROCESS AND CHORE MAGE:4155401 5	602019/7011 NOL COST CHICA TO THE COST CHICAGO STATES COST CHICAGO DVANDAI MBNA martial cost	Arabidopsis utalialia 1937 2-1 ili. 1937 Jane 4 Juneau hana elalonotelin (BNSP) dene, exons 2, 3 and 4	PC3 CN00A2,310800-024-d11 GN0042 Homo sapiens cDNA	Social money remarks the suppressor of MIF2 Sm(4p (SMT4) gene, complete cds	Saccington 3000 CGAP Bi2 Homo saptens cDNA clone IMAGE:1632465 3' similar to WP:C37A2.2	CE08011;	oue3b05.s1 NCI_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1632465 5 similar to Wr. Co. A2.2	CE08611;	0059d02/y5 NCL_CGAP_Lu3 ndiilo sapidis construction construction alamant i	oc59402 y5 NCI_CGAP_Lu5 Home saplens cDNA clone IMAGE:1570467 5' similar to contains L1.t3 L1	repetitive element;
Single Lyon Figure	Top Hit Database Source		HOMAN		EST HUMAN	Г							LN LN			HUMAN		EST HUMAN	Τ			L _N		EST HUMAN	Ę.	12	NAMOE 101	Z	EST HUMAN		EST HUMAN		EST HUMAN	EST HUMAN
fille	Top Hit Acession No.		.1		17.1						,	,	02 AF217796.1	02 AJ005375.1	4503034 NT	02 BE250008 1	3EZ 30000. I	00 00 00 00	11005053.1		.02 AF030694.2	6681044 NT	6681044 NT	-02 BF348454.1	-02 AB008019.1	-02 L 24757.1	-02 BF368016.1	7.9E-02 U27832.1	7 OE 02 A1081644 1		7.9E-02 A1081644.1		-02 AI793275.1	-02 AI793275.1
	Most Similar (Top) Hit BLAST E	Value	8.0E-02 AI434202.1	8.0E-02 X72794.1	8 OF-02 A	8 OF 02 4	7 20 20 0	0.0E-04	0.0E-02/AL11994	8.05-02/	8.0E-02 X/4208.1	8.05-02/	8 OF -02	8.0E-02	8 0F-02	7 05 03	70-28: /	100	1.85-02		7.9E-02	7.9E-02	7.9E-02	7.9€	7.9€	7.9E	7.9E		7 01.03	1.95-04			7.85	7.8
	Expression Signal		1.43	6.33	78.0	0.00	3.10	1.82	3./8	1.12	1.12	0.55	700	939	224	1 0	3.52		67.7		0.97	5.01	5.01				1.16	2.79		4.21	4.21		1.77	1.77
	ORF SEQ ID NO:		30093		67,700	30443	31516	31518	١		35128		00000				27634		28392		29168			30011		30204		33754		5 35773	35774		0 26604	26605
	Exon SEQ ID		17457	47.40	200	ı	- 1		20722		21956	22706		23362		\perp	14900		15744		16529		1_	L	1_	17581	19360	1_		7 22575	7 22575	_	8 13940	13940
	Probe SEO ID	<u>.</u>	17.75	201	4/04	5108		7080	8027	9289	9289	10058		10692	C8171	12/48	2171		2978		7777	3832	2823	4645	4760	4851	629)	7931		9927	0027		1188	1188

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Homo saplens zinc finger protein 92 (ZFP92), expressed-Xq28STS protein (XQ28ORF), and biglycan (BGN) genes, complete cds; and plasma membrane calcium ATPase isoform 3 (PMCA3) gene, partial ods Homo sapiens zinc finger protein 92 (ZFP92), expressed-Xq28STS protein (XQ28ORF), and biglycan (BGN) Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1b mRNA, complete Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1b mRNA, complete ta80b08.x1 NCL_CGAP_HSC2 Homo saplens cDNA clone IMAGE:2050359 3' similar to gb;Z28876 60S Homo sapiens ASCL3 gene, CEGP1 gene, C11orf14 gene, C11orf15 gene, C11orf16 gene and C11orf17 te80b08.x1 NCI_CGAP_HSC2 Homo septens cDNA clone IMAGE:2050359 3' similar to gb:226876 60S RIBOSOMAL PROTEIN L38 (HUMAN); genes, complete cds; and plasma membrane calcium ATPase isoform 3 (PMCA3) gene, partial ods Homo septens partel AF4 gene, exons 2 to 7 and Alu repeat elements
Gallus gallus collagen type XII alpha-1 (COL1241) gene, promoter region and partial cds
zu53d11.r1 Soares overy tumor NbHOT Homo saplens cDNA clone IMAGE:741717 5' similar to
TR:G1173905 G1173905 SPLICEOSOME ASSOCIA TED PROTEIN; Homo saptens interferon regulatory factor 7 (IRF7), mRNA 601316428F1 NIH_MGC_8 Homo saptens cDNA clone IMAGE:3634903 6' EST112214 Cerebellum II Homo saptens cDNA 6' end similar to similar to protocadherin 43 HIV-1 strain 97USNG30 from USA, envelope glycoprotein (env) gene, partial cds no88b06.r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:771731 Bacillus subtilis complete genome (section 21 of 21); from 3889281 to 4214814 an25g02.x1 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:1699730 3' 601440439F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3925449 67 S. S. cerevisiae CAT8 gene PROBABLE SERINE/THREONINE-PROTEIN KINASE YBR059C Top Hit Descriptor Human interleukin-11 receptor alpha chain gene, complete cds Homo sapiens KIAA0628 gene product (KIAA0628), mRNA Homo sapiens WRN (WRN) gene, complete cds RIBOSOMAL PROTEIN L38 (HUMAN); Single Exon Probes Expressed in Brain 8 EST_HUMAN Top Hit Database Source HUMAN EST HUMAN HUMAN **EST_HUMAN** EST_HUMAN EST HUMAN SWISSPROT EST_HUMAN EST HUMAN EST. Ę Ę Ę Top Hit Acession 11422757 114368 7.8E-02 U32323.1 7.8E-02 AF096349.1 7.7E-02 AF181897.1 7.7E-02 AJ238093.1 X78344.1 7.8E-02 AA469354.1 7.8E-02 Z99124.1 7.8E-02|BE250048 ġ 7.8E-02 AF233437.1 7.8E-02 AF233437.1 7.7E-02 AA402949.1 7.7E-02 P38080 .6E-02 AA298447.1 BE514432.1 7.8E-02|U82695.2 7.8E-02 U82695.2 7.7E-02 AI318662.1 7.6E-02 AJ400877.1 7.6E-02 AI061275.1 7.7E-02 Al318662.1 7.7E-02 114 7.8E-02 7 (Top) Hit BLAST E Vælue Aost Similar 7.6E-02 7.7E-02 0.88 0.71 Expression. Signal 0.88 99.0 0.79 0.62 4.58 3.92 5.37 3.94 0.85 4.51 1.97 0.71 0.81 ORF SEQ ID NO: 32479 32480 34520 34618 34789 35183 35547 38494 34790 31145 26800 33625 35590 35895 31749 35896 36863 28798 28812 SEQ ID 16484 19457 19457 21642 21471 21642 20504 16329 18255 22386 23613 22879 16302 16141 18161 18787 Probe SEQ ID 8978 8976 5027 8779 8951 7809 0562 12754 8951 ö 9261 3574 5456 9735 10933 3382 3403 10031 10031 3547 9009

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	Top Hit Descriptor	601236402F1 NIH_MGC_44 Hamo sapiens cDNA clane IMAGE:3608401 6'	Homo saplens SCL gene locus	Campylobacter ieluni NCT C11168 complete genome; segment 5/6	DC1-HT0545-020800-017-d06 HT0545 Homo sapiens cDNA	NOTIFICATION OF THE MACE RT Home seriens cDNA clone IMAGE:3839810 3'	00100481901 IVIII_INOC_C IVIII	Lesculantini milyta to trose phosphate translocator	L'esculentum minara de mode propries con series con Anno Serie		Homo sapiens solute carrier family 6 (neurotransmitter transporter, glycine), member 9 (SLC6A9), mRNA	Homo saplens solute carrier family 6 (neurotransmitter transporter, glycine), member 9 (SLC6A9), mRNA	Homo sapiens IL-18 gene for interfeutrin-19, introd I have axin 2	wq24h09x1 NC_CGAP_KId11 Homo sapiens CUNA GOTE INACCESSES AS SINGLES IN THE WINDOWS AS SINGLES I	WISSEQ X1 NCI CGAP Brn25 Home septens cDNA clone IMAGE 2426491 5 SITILIES CON SECTION OF THE PROPERTY OF THE P	ENOLASE (HUMAN); A 141-6013 HEMBA Homo seniens cDNA clone HEMBA1000264 5'	ACTION OF THE PART HOME SEPTENCE CON COMPANY SIGNIFICATION STATES AND CONTRINS SEPTENCE OF THE PART OF	MER27 repetitive element: MER27 repetitive element: ACT repetitive el	010/02/03F Militaria (1977)	C:IIIII DSIM 29 113 133 133 134	Fruine hemesyins 4 strain NS80567, complete genome	Admis miscrilis paired-like homeodomain transcription factor 1 (Pitx1), mRNA	witarhot vt Soares NPL T GBC S1 Homo saplens cDNA clone IMAGE:2358385 3'	Home series ADP/ATP cerrier protein (ANT-2) gene, complete cds	Dotting Sprains Activity receptor like kinase 1 (Acvit), mRNA	National State of Sta	Mississocial Progress infant brain 1/1/18 Homo sapiens cDNA clone IMAGE:32339 5	191-1920. To accommend the septent CDNA clone IMAGE:3895284 5	60 (1453300) 1 (11) 1 (10) 2 (10) 1 (hazara va NCI CGAP GU1 Homo sepiens CDNA clone IMAGE.2967861 5' similar to SW SCA2, HUMAN	015127 SECRETORY CARRIER-ASSOCIATED MEMBRANE PROTEIN 2.;
Single Liver of the color	Top Hit Database Source	EST HUMAN 6	Γ		THE PERSON	Т	HOMAN		Т	ESI HOMAIN			TN	EST_HUMAN	Г	\neg	EST HOMAIN	HUMAN	HOMAN	П	- HOME		HAAM	NAMOL -			14444111	Ţ	HOMAN	Ž	EST_HUMAN
Builo	Top Hit Acession No.	T			T	1	32			02 AW 996645.1	5902093 NT	5902093 NT		7.5E-02 Al948714.1		02 Al864367.1	02 AU116913.1	-02 BF221730.1	-02 BF206809.1	-02 X79460.1	7.4E-02 AW838547.1	7.4E-02 AF030027.1	1N 690cc/9	7.4E-02 AI807885.1	7.4E-02 L78810.1		6678492 N I	7.4E-02 R17477.1	7.4E-02 BE880112.1	7.4E-02 U56089.1	7.4E-02 AW629605.1
	Most Similar (Top) Hit TBLASTE Value	7 RF-02 RF379328.1	100 101	νIσ	7	7.6E-02 B	7.6E-02 B	7.6E-02 X92656.1	7.6E-02	7.6E-02	7.5E-02	7.5E-02	7.5E-02/	7.5E-021/		7.5E-02	7.5E-02	7.5E-02	7.5E-02	7.5E-02	7.4E-02	7.4E-02	7.4E-02	7.4E-02	7.4E-02	7.4E-02					
ŀ	Expression Signal	8	092	1.4/	1.83	0.49	0.75	0.71	0.71	245	1.44	1.44	1.17	160	2	1.05	1.17	0.54	7.0	0.72	1.46	0.92	1.32					2.18		1.03	1.12
	ORF SEQ ID NO:	0,000	32012	35194		35979		36359		37595	26199	28200		1		34066			36252		25891	2			30024			2	33618	34228	3 34913
	Exon SEQ ID NO:	1	19037	22024	22448	22767	22895	23133	23133	24273	١	<u> </u>	-1_	1	1835	20830		22580	L	L	13250	14192	15299	16336	Ŀ	17473	17616	19172	20496	21092	4 21753
	Probe SEQ ID NO:		6263	9270	9797	10119	10247	10487	10487	11678	767	707		7/55	29/62	8236	8405	9932	10390	10488	465	1445	2585	3581	4656	4741	4889	6403	7801	8389	9064

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					S	ngle Exon Pr	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Simller (Top) Hit BLAST E Value	Top Hit Acession	Top Hit Database Source	Top Hit Descriptor
9064		34914	1.12	7.4臣	-02 AW629605.1	EST HIMAN	hh67d11.y1 NCI_CGAP_GU1 Homo septens cDNA clone IMAGE:2967861 5' similar to SW:SCA2_HUMAN
9336			0.52	7.4	-02 A 167 2939 1	FST HIMANI	WEZAMO W SOUTH CARRIER-ASSOCIATED MEMBRANE PROTEIN 2.
8339			0.52	7.4E-02	7.4E-02 AI672939.1	EST HUMAN	we74402 v1 Sparse Disclorance Colon INHCD Homo sepiens cDNA clone IMAGE:2346819 31
9714	\Box		1.03	7.45-02	7.4E-02/U62293.1	LN	Human I Mukingara and all the first the section of
9841		35692	0.52	7.4E-02	7.4E-02 BF512678.1	EST HIMAN	The BMT own of the William of the West Spliced LIM-Anasel (LIMK1) gene, complete cds
10939	_	36869	1.26	7.4E-02	7.4E-02/AA059167.1	EST HIMAN	764401 1 Source cells NO. COAP. Sub7 Homo seplens cDNA clone IMAGE:3069898 31
12126	- 1		1.53	7.4E-02	11525893	LV	Homo canions blotter of
12381			2.21	7.4E-02	AW379	EST HIMAN	CMALHTO2/2 DB4200 027 111 1120 027 111 111 1120 027 111 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110 027 1110
456			1.5	7.3E-02	02 BE9649612	EST HIMAN	COLDEREZZODA NILL ACCOUNTY AND Saplens cDNA
458			1.5	7.3E-02	02 BE964961 2	EST HIMAN	BOT GOAD SONT INITIA WISC. 69 Homo saplens cDNA clone IMAGE:3886209 3'
699	13445	26085	3.9	7.3F-02	02 AF001780 1	TIONOL TIN	OU DOO/ SONT INIT MICC. 69 Homo septens cDNA clone IMAGE:3886209 3/
1464	LI	26900	3.62	7.3E-02		EST LIMAN	I nermotoga maritima section 101 of 136 of the complete genome
1837	15580		12.41	7.3E-02		EG - HOWAN	CWC-NN 1004-130300-284-g08 NN 1004 Homo sapiens cDNA
				1	1	Z	Homo saptens chromosome 21 segment HS21C102
6361		32126	1.32	7.3E-02	7.3E-02 AA779977.1	FST HUMAN	224a02.s1 Soares fetal_liver_spleen_1NFLS_S1 Homo sapiene cDNA clone IMAGE:451178 3' similar to
7368	20048	33128	2.58	7.3E-02	l	SWISSPROT	PROTECTION TO TAKE SOUTH 4 (HUMAN);
7368	20048	33129	2.58	7.3E-02		SMISSEDDA	POLICE TO THE PROPERTY OF THE
8008	20762		1.15	7 3E 03	7 3E 02	- CALLOS	
9110	21798		1 14	7 3E 02		Z I	Home sapiens KIAA0424 protein (KIAA0424), mRNA
				1.35.02	ADD 1 1090.1	Z	Homo sapiens mRNA for KIAA0518 protein, partial cds
11179	19131	32126	2.06	7.3E-02	AA779977.1	EST HUMAN	224802.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:451178 3' similar to
11844	24428		5.07	7.3E-02	7.3E-02 11560138 NT	E	Reftus norvegicus caspase recruitment domain protein of 100 24434
117	12837	25577	1	7.2E-02	7.2E-02 AE000882.1	F	Methanobacterium thermoautotrophicum from bases 1029155 to 1039934 (section 88 of 148) of the complete
117	12837	25578	-	7 25.03	7 2E-02 AE000882 4		Methanobacterium thermoautotrophicum from bases 1029155 to 1039934 (section 88 of 148) of the normalists
1458	14205	26890	2.72	7.2F-02	T	LIV.	gename
1458	14205	26891	272	7 25 02	7 25.02 41 463304 2	1	norno sapiens chromosome 21 segment HS21C101
			1	1.45.7		Z	Homo sapiens chromosome 21 segment HS21C101
2852	15267	1	2.83	7.2E-02	U14794.1	Ę	numunodenciency virus type 1 isolate 26 reverse transcriptase (pol) gene, internal fragment, partial cds
3863	16615	29254	0.95	7.2E-02	AW 298322.1	EST HUMAN	ULH-BWO-ail-a-05-03-11 st NICL COAP St-be Hammer
4312	17051	29678	4.65	7.2E-02	BF572307.1	Т	60207775771 NIH MGC R2 Home received and supplies CDNA clone IMAGE:2732049 3'
4844	17378	30010	0.7		36563		Rhodomonas salina mitochondrion, complete ganome
							Allow Actions to the second se

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Single Exon Probes Expressed in Diam.	Top Hit Descriptor	Methanococcus jannaschii section 73 of 150 of the complete genome	CALMODULIN	601883558F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4095710 5	Streptococcus pneumoniae putative response regulator (zmpR), putative histidine kinase (zmpS), and putative	zinc metalloprotease (zmpB) genes, complete cds	Strongylocentrotus purpuratus mitochondrion, compiete genome	PROLINE-RICH PROJECTIVE AND 2	PROLINE-RICH PROLEIN WITS	Lactococcus takens capit games hadisa diabulin (SHRQ)	Human gene rot sex normane-bulloung globbullin (cr. 15.0)	AV712452 D.C.A. Homo sapiraris curis D.C.A. Social Control of the Article products,	Homo sapiens plasma membrane cardum A i rass isolomi i (A i . 227, 550.5, 200.5) partial cds	AN 1783 F 23 F 1 NIH MGC 20 Homo sapiens cDNA clone IMAGE: 4026436 5	2017 Control Control Control Home saniens CONA clone (MAGE:3120333 3' similar to TR:Q9Z340 Q9Z340	ATYPICAL PKC SPECIFIC BINDING PROTEIN;	os62c07.s1 NCI_CGAPGCB1 Homo sapiens curve curve in notice in the control of th	(BGN)	Homo sapiens zinc finger protein 92 (ZFP92), expressed-Aqzes is protein (Azzootti, , and beywardense complete cds; and plasma membrane calcium ATPase isoform 3 (PMCA3) gene, partial cds	2012/2/2026 NIH MGC 53 Homo saplens cDNA clone IMAGE:3685951 5	6010E810AF1 NIH MGC 10 Homo sapiens cDNA done IMAGE:3451559 5'	328-05 s.1 Shares, felal liver spleen 1NFLS S1 Homo saplens cDNA clone IMAGE:451641 3	Define noneclars hell Hitranscription factor Mist1 (Mist1) gene, complete cds	name ambas in region franchise transmembrane protein dectin-1 mRNA, complete cds	And Andreas NhHMPu S1 Homo sapiens cDNA clone IMAGE:1048398 5	BIO 1804-1, 1 COMES Anim III Commission (P) Homo sepiens cDNA clone PS 13D5 3'	Autoria por información de la TM) dene. complete cds	Horing segment agreement and a company of the compa	CMM-NN I Judg-Zubosud-1 I Grott I Military 2010 1 Company I Judg-Zubosud-1 I Grott I Military 2010 1 Company I Judg-Zubosud-1 Company I Judg-Zubos		Human immunodeficiency virus type 1 (D9) provital structural capsid protein (gag) gene, partial cds	6018/2281F1 NIH MCC 33 Fruit Barriers CDNA clone IMAGE:1736922 3	No. a Company of the
le Exon Probes	Top Hit Database Source	N-	SWISSPROT C	Т	Τ			┑	ISSPROT		T	EST HUMAN A	I à	THE PARTY	EST HUMAN	EST_HUMAN A	EST_HUMAN @		<u> </u>	14494111	Т	ES HOMONIA	Т		т	EST HUMAN	Т	т	EST HUMAN	┰		- 1	LEST_HUMAN
Sing	Top Hit Acession No.	22 U67531.1	10 044420	12 RE216086 1		02 AF221126.1	5834897 NT	-02 P05143	P05143	02 Y17217.1	02 X16349.1	02 AV712452.1	1 4 4564 4	L14301.1	-02 BF125399.1	02 AW873187.1	-02 AA768204.1		0	-02 082695.2	-02 BE565003.1	-02 BE539214.1	7.2E-02 AA706897.1	7.2E-02 AF049874.1	7.2E-02 AY009090.1	2 AA773696.1	7.2E-02 AJ230796.1	7.2E-02 U82828.1	7.2E-02 AW900962.1	-02 AF020439.1	7.1E-02 L02290.1	7.1E-02 BF208802.1	7.1E-02 AI125264.1
	Most Similar (Top) Hit BLAST E Value	7.2E-02	7 25 03	7 25 02	1,45.72	7.2E-02	7.2E-02	7.2E-02	7.2E-02	7.2E-02	7.2E-02	7.2E-02		1.45-02	7.2E-02	7.2E-02	7.2E-02		1	7.2E-02	7.25-02									7.2			
	Expression Signal	2,88		10.1	1.30	0.64	1.5	8.0	0.8	190	0.49	2.32		3.8	0.93	2.53											4.45	1.73	8.19	3.62	2.01		0.77
	ORF SEQ ID NO:	30635	00000	30636	32824	32840	L	33910		L		35346			35667	35747				36108	36234				37775	31113				2	27344	L	33622
	SEQ ID	- 1	-1	- 1	19759	40775		1.	20781	L	L	1.	١.	22311	22465	22552	┸			22898	23018	5 23041	23155	23512	ı	24560	24583	L	1_		7 14634	1	
	Probe SEQ ID NO:	3002	cnze	5206	2088	7002	7 083	2887	8087	8967	9474	9511		9659	9814	800	3	10092		10250	10372	10395	10509	10830	11849	12035	12069	12182	12196	12599	1897	238	7807

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Table 4
Single Exon Probes Expressed

Single Exon Probes Expressed in Brain	Top Hit Descriptor	143074E1 NIEL 1600 15 II	COLLAGEN ALDUA 4/X4/X CLANDER SEPTEMBER CON CONTRACTOR STATEMBER CONTRAC	Marfallo Micrit 1 cons	Stody of Strategies of the stody of the state of the stody of the state of the stody of the state of the stat	200104-81 Strategene colon (#837204) Homo sapiens cDNA clone IMAGE:609599 3/ UI-H-BIT-ecy-c-07-0-UI st NCI CGAP Silika Docestics Child	at65a12.s1 Soares_tests_NHT Horno capiens cDNA clone 1375878 3' similar to gb:K03002 60S	CMAN-I MONOGO 270 - 40 I II 1000 I I	Canis familiaris individual mitto add a mitto a mi	601816291F1 NIH MGC 58 Home control of the control	Lumbricus rubeilus mRNA for cucloristilla B	AV689285 GKC Homo sabians chiva choca access	Gallus gallus mBNA for partial aczonin YI entired invited for	African swine fever virus, complete nanome	Rat to nermine encilon H. Abain near Brown at an	Human myosin binding brotein H (MARP-H) page commissions	ah99a05.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1327184 3' similar to gb:L14837	TOTAL SONOTION PROTEIN ZOTA (HUMAN);	Homo sapiens chromosome 21 segment HS21C010		rigino sapiens regulator of Gz-selective protein signaling (ZGAP1) mRNA, and translated products	28S PROTEASOME REGILIATORY SUBUNITIES (NUCLEAR ANTIGEN 2107)	Entercoccus faecium cystehe aminoseptidase (nenc) gene media de nence de la company de	(bgiB), beta-glucoside specific transport protein (bgiS), transcription antiterminator (bgiR), entercein R	precursor (entB), entercoin B immunity prote>	ou i rezassi i Nid MGC 7 Home sapiens cDNA clone iMAGE:3636253 6	Canine distember virus strain A75/17, complete genome	Fruman carmodulin (CALM1) gene, exons 2,3,4,6 and 8, and complete cds	601340681F1 NIH MGC 53 Home capiens CDNA close IMAGE:3883030 5'	Barbarie duck parwumis REP modein (can) and the control of the con
gle Exon Prot	Top Hit Database Source	EST HIMAN	SWISSPROT	Ę	EST HIMAN	EST HUMAN	Г		Т	Т	Į.	Т	Т		Ė		TO HOLD	T			TORDEO!	Т	Т		TOL	Т		T HUMAN	Т	N
Sin	Top Hit Acessian No.	7.1E-02 BE304784.1	007092	7.0E-02 X96677.1	7.0E-02 AA056343.1	7.0E-02 AW138152.1	-02 AA815438.1		T	7.0E-02 BF381987.1	ŀ	-		9628113 NT	02 K02901.1		12/44724205-1		Γ	2002	4307.800	02 Q06364				T	T		1	
	Most Similar (Top) Hit BLASTE Value	7.1E-02	7.0E-02/Q07092	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02	7.0E-02 Y19187.1	7.0E-02	7.0E-02	7.0E-02	7.0F-02	8 9F-02 A	6.9E-02	20 0	6.9E-02/C	6.9E-02		- CO	R OE. 02 DE 28480E 4	6 9E-02 A E164067 4	6.9E-02 112022 1	6.9E-02 BI	6.9E-02 BE567435.1	6.9E-02 U22967.1
	Expression Signel	6.41	-	1.27	1.08	2.1	0.74	1.28	1.06	7.24	0.57	1.29	0.84	1.26	1.24	0.51	4.98	4.3	4.3	1.0	1.41	1.41		Č	1 25	0.61	1.12	1.01	1.01	0.7
	ORF SEQ ID NO:		25931		27199	28440	29268		29560	30227		33059	33271	34836	35331	36702	37338	25917	25918		29163	29164		30448	30462			34282	34283	34860
	Exon SEQ ID NO:	24483	13299	14233	14498	15783	16628	16861	16930	17604	18088	19983	20177	21686	22150	22502	24035	13285	13285	14058	16525	16525	-	17831	17845	20187	20646	21143	21143	21708
ſ	Probe SEQ ID NO:	11922	515	1486	1756	3027	3878	4119	4189	4877	5283	99	7508	8996	9497	8827	11345	501	501	1310	3773	3773		5113	5127	7516	7951	8451	8451	8018

Page 131 of 536 Table 4 Single Exon Probes Expressed In Brain

					ino ino	אום דייטון דייטג	Single Exoli Plobes Expressed in Brain
Probe SEQ ID S NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
12232	24685		1.69	6.9E-02	P44621	SWISSPROT	PROTEIN TRANSPORT PROTEIN HOFC HOMOLOG
12447	24817		1.46	6.9E-02	AF195953.1	TN	Homo sapiens membrane-bound aminopeptidase P (XNPEP2) gene, complete cds
1875	14613	27321	1.56	6.8E-02	AA496759.1	EST_HUMAN	ae30f02.rl Gessler Wilms tumor Homo saplens cDNA clone IMAGE:897339 5' similar to gb:M22382 MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR (HUMAN);
1875	14613	27322	1.56	6.8E-02	AA496759.1	EST_HUMAN	ae30f02.rl Gessler Wilms tumor Homo saplens cDNA clone IMAGE:897339 5' similar to gb:M22382 MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR (HUMAN);
1900	14637	27346	3.77			LN	Homo saplens putative hepatic transcription factor (WBSCR14) gene, complete cds
3097	15862	28503	1.19	6.8E-02		EST_HUMAN	al75a06.s1 Soares_testis_NHT Homo sapiens cDNA clone 1376626 3
3097	15862	28504	1.19	6.8E-02	AA781996.1	EST_HUMAN	ai75a06.s1 Soares_testis_NHT Homo sapiens cDNA clone 1376626 3'
3097	15862	28505	1.19	6.8E-02	AA781996.1	EST_HUMAN	ai75a06.s1 Soares_testis_NHT Homo sapiens cDNA clone 13766263'
4516	17251		0.86		BE141076.1	EST_HUMAN	MR0-HT0069-071099-001-c05 HT0069 Homo saplens cDNA
6525	19291		9.0	6.8E-02		SWISSPROT	CELL-SURFACE RECEPTOR DAF-1 PRECURSOR
623	19460		1.09		BE061890.1	EST_HUMAN	RC1-BT0254-090300-017-d09 BT0254 Homo sapiens cDNA
7180	19866	32939	8.73		.2	NT	Homo saplens chromosome 21 segment HS21C068
7584	20252	33358	0.63	6.8E-02		NT	Dictyostellum discoldeum myosin heavy chain kinase A (MHCK A) mRNA, complete cds
8186	20880	34017	5.01	6.8E-02		NT	Pyrococcus abyssi complete genome; segment 5/6
8186	20880	34018	5.01	6.8E-02	AJ248287.1	NT	Pyrococcus abyssl complete genome; segment 5/6
11873	25379		23	6.8E-02		EST_HUMAN	FB4A8 Fetal train, Stratagene Homo sapiens cDNA clone FB4A8 3'end similar to LINE-1
12001	24537		2.85	6.8E-02	AA758014.1	EST_HUMAN	ah67f05.s1 Soares_testis_NHT Homo sapiens oDNA clone 1320705 3'
12551	24888		1.65	6.8E-02	AW975839.1	EST_HUMAN	EST387948 MAGE resequences, MAGN Homo sapiens cDNA
12613	24920		3.06	6.8E-02	10585	LN	Mus musculus latent TGF beta binding protein (Tgfb), mRNA
1519	14266		1.93	6.7E-02	AF115536.1	NT	Oncorhynchus mykiss TAP1 protein (OnmyTAP1) mRNA, OnmyTAP1*01 allele, complete cds
1886	14623	27333	2.27	6.7E-02	AI220285.1	EST HUMAN	qg79e04.x1 Soares_NFL_T_GBC_S1 Home saplens cDNA clone IMAGE:1841406 3'
3706	16459	28097	4.52	6.7E-02	P17278	SWISSPROT	HOMEOBOX PROTEIN HOX-D4 (CHOX-A)
7749	20445	33567	0.55	6.7E-02	X62695.1	LN	H.sapiens DNA for cGMP phosphodiesterase (exons 4-22)
7749	20445	33568	0.55	6.7E-02	X62695.1	ΤN	H.sapiens DNA for cGMP phosphodiesterase (exons 4-22)
	21030	34167	0.47		AW082688.1	EST_HUMAN	xb61c11.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2580788 3'
	22153	35333	0.69	6.7E-02/	AW137359.1	EST_HUMAN	UI-H-BI1-acr-g-01-0-UI s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2715433 31
9500	22153	35334	0.69	6.7E-02	AW137359.1	EST_HUMAN	UI-H-BI1-acr-g-01-0-UI.s1 NCI_CGAP_Sub3 Homo sepiens cDNA clone IMAGE:2715433 3'
1348	14096	26771	1.07	6.6E-02	AF245116.1	NT	Drosophila mèlanogaster cactin mRNA, complete cds
2180	14909	27641	3.31	6.6E-02	1.1	Z	Mus musculus Capn12 gene for calpain 12, exons 1-21, three alternative transcipts
3456	16212	28865	10.57		R64306.1	EST_HUMAN	y18b10.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:139579 3'
3471	16227	28881	2.59	6.6E-02	7108357 NT	NT	Homo sapiens mesothelin (MSLN), transcript variant 1, mRNA
3471	16227	28882	2.59	6.6E-02	7108357 NT	Ę	Homo saplens mesothelin (MSLN), transcript variant 1, mRNA

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens TESTIN 2 and TESTIN 3	INTER-AL PHA-TRYPSIN INHIBITOD DEALY CONDICE das, alternatively spliced	INTERAL PHA-TRYPSIN INHIBITOR DEALY CHAIN HE PRECURSOR (I'II HEAVY CHAIN HE)	P. Autoris mRNA for chalcane symptoms	MATERNAL PERECT PROTEIN STATISEN	MATERNAL FEFE OF DROTTEN STATISTIC	Homo sapiens chemokine recentor CXCR4 asse commuter and a second of the	Dictivistely discorder my define dark years again to be a complete day	DNA POLYMERASE ZETA CATALYTICS I IBINIT / LIDES (2)	Human respiratory syncytial virus complete agency	Human respiratory syncytial virus complete gendine	197a06.XT NCI CGAP Lit24 Home seniors all A characters at 127.22.22.22.22.22.22.22.22.22.22.22.22.2	Homo sepiens FWS nar22 mod bom 22 and bom 22 mod bom 22	Homo saplens when lin (VC) I m BNA	MR1-SN0084-0108nn-nn6-a12 Sninga Linne and and and and and and and and and and	Homo saplens chemokine recentor CYCD4 and	Mus musculus DIPR gene / Diph https://www.new.complete.cds	Refits bonedicies constrained Dies 254 (Oversta	601671046F1 NIH MGC 20 Home soules all Latter 1, 140 CH 2000 F1	Homo sapiens E2F-like protein (1 OCK127n) with a manual contraction of the contraction of	Xenopus laevis alpha/E catenin mRNA complete cate	Aquifex aeolicus section 96 of 109 of the complete canno	2/46/h12.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:788743 3' similar to gb:M28038	Azotobacter vinelandii ATCC 9046 nagatiya azoti later M BY BY	301656817R1 NIH MGC 67 Home emiliar cDNA along MACE access 6	601856817R1 NIH MGC 67 Home septems clove clone imAdE;3865637 3	601823511F1 NIH MGC 77 Home saplens CDNA clone IMAGE:3066637 3	2/3/2905.s1 Sogres NhHMPu S1 Homo saplens CDNA Appa NA CE GERALL 21	Rabbit microsomal epoxide hydrolese	Nectria haematococca kinesin related protein 2 (KRP2) some committee 3.3.	A carterae precursor of peridinin-chlorophylla-poylen (DOD)	Mus musculus histone deacetylase 5 (Hdac5) mRNA	Mus musculus histone deacetylase 5 (Hdac5), mRNA
gle Exon Pro	Top Hit Database Source	TN	SWISSPROT	SWISSPROT	LZ	SWISSPROT	SWISSPROT	LN	NT	SWISSPROT	Ľ	LZ	EST HUMAN	LZ	5	EST HUMAN	N	5	LN	EST HUMAN	ート			$\overline{}$	L	т	Т	Т	HUMAN	7				
Sin	Top Hit Acession No.	6.6E-02 AF280225.1		6.6E-02 Q61703				72.1	-02 AF006055.1		9629198 NT	9629198 NT	02 AI458752.1	02 Y07848.1	1430559	02 BF374248.1		37891	02 AF167430 1	Γ	99096	02 U47624.1	7.	6.6E-02 AA443991.1	-	2	Π				6.5E-02 AF102993.1 N		6996923 NT	6996923 NT
	Most Similar (Top) Hit BLAST E Value	6.6E-02	8.6E-02	6.6E-02	8.6E-02	8.6E-02	6.6E-02	6.6E-02	6.6E-02	8.6E-02	6.6E-02	6.6E-02	8.6E-02	8.8E-02	8.6E-02	6.6E-02	6.6E-02	6.6E-02		8.6E-02	6.5E-02	6.5E-02 €	6.5E-02	6.6E-02	6.5E-02(6.5E-02	8.5E-02	8.5E-02 E	6.5E-02 A	6.5E-02 M21496.1	6.5E-02 A	6.4E-02 X94549.1	8.4E-02	6.4E-02
	Expression Signal	1.29	7.03	7.03	3.44	0.58	0.58	1.81	0.84	0.63	0.58	0.58	0.65	1.66	0.63	6.88	1.46	2.66	1.38	2.49	1.32	3.08	1.77	2.03	0.95	0.55	0.55	0.48	6.56	3.73	4.66	2.09	0.96	1.18
	ORF SEQ ID NO:	29436	30261		32258	32286	32287	33870	34208		34654	34655	35701	35845		36811	37793			25977	26398	26793	27169	31162	30549	35693	35694	36225	36466			25970	28429	28429
	Exan SEQ ID NO:	16805	17649			19283	1	ı			. 1		- 1	ı	22670	23563	24451	24812	25006	13349	13732	14118	14470	18270	17953	22493	22493	23010	23232	24463	24691	13343	15780	15780
	Probe SEQ ID NO:	4060	4921	4921	6489	6701	6701	7847	8372	8678	88 19	8819	9851	8884	10022	10883	11867	12442	12740	268	88	1370	1728	5471	6877	9842	9842	10363	10635	11894	12240	561	3014	4839

Page 133 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	qe07b01.x1 Soares_bestis_NHT Homo sapiens cUNA cione IMACE:1730249 3 Sillilia & Contain Contai	Mus musculus IFN-response element puruling racking (MC-eng-1) gene, complete cds	Helerodera glydnes bera-1, 4-andogludariase-1 produsor (NG-eng-1) gene, complete ods	Heterodera grydnes beta-1, -reinogrowango i promissione i promissione i MACE: 2346790 31	Wel 3g LZA1 Suggest Orchig 2012 St Home saplens cDNA clone IMAGE:3950503 31	Nelsearla meninnilidis seroaroup A strain Z2491 complete genome; segment 6/7	Mais miscailus chanegrann subunit 6a (zeta) (Octoa), mRNA	witch incommon one-process that is ambda ZAP Express Homo saplens cDNA 5	NITH CONTROL HIMMON MENUAL FORM CIGAR stem cells Home sapiens cDNA clone CBDAIA10	AFTOCIONAL ALGORO A44 CORON Home saplens cDNA	RCI-C LOSSY-19000-01-1900 C LOSSY INCHES COS	Homo saprens miny of its Novembers of the (TOP28) gene, expns 16, 17, and 18	Homo saptens DNA tongloomeree II beta (TOP2R) cene. excus 16, 17, and 18	חום פשטופיוז ביוער וסף מפתיום מספר וו בכים (יייי ביייי) פייייי	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	111-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	Homo sapiens mucin or Innovative personal sapients and the control of the control	נוספטשוום ווומאסים וווו איני וכן יוסאיים איני ווויסאיים איני וווויסאיים אווויסאיים אווויסאיים איני וווויסאיים איני ווויסאיים איני ווויסאיים איני ווויסאיים איני וווויסאיים איני ווויסאיים איני ווויסא	Mus musculus major histocompatibility locus ciass ili regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, CLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes	HEAT SHOCK PROTEIN 70 HOMOLOG	601873316F1 NIH_MGC_54 Hamo sapiens cDivA clude invace	H. sapiens gene encoding La autoantigen	Drosophila metanogaster Dollilla gene, exclisi 100	Heparus G Vilus RYA IOI POYACETTI (CONTRACTOR OF AVAGABATA CAKE Home sapiens CDNA clone GKCAHE01 5	601873316F1 NIH MGC 64 Homo sapiens cDNA clone IMAGE: 4097499 5'	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 68	
Single Exoll Flobes Explored	Top Hit Database Source	est_HUMAN LT			T	Т	HOMAN			T	Т	EST HUMAN R	7		E L	H)				Į.	F Z		EST_HUMAN 6	П		7	-	FIGURE FIN	
Buic	Top Hit Acession No.	6.4E-02 A1191956.1	7305186 NT		6.4E-02 AF052733.1		6.4E-02 BE974448.1		6753323 NI	6.4E-02 AA093305.1	6.4E-02 AF150195.1		1	02 AF087150.1	02 AF087150.1	02 U91328.1		02 091328.1	AF107890.1	6.4E-02 AJ277174.1	02 AF109805.1	-02 P37092	-02 BF210736.1	.02 X97869.1	-02 AJ243916.1	6.3E-02 AB010162.1	-02 AV698070.1	6.3E-02 BF210736.1	ZJAL1013/4.4
	Most Similar (Top) Hit BLAST E Value	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.4E-02		6.4E-02	6.4E-02	6.4E-0;	9.3E	6.3E	6.3E	6.3E		6.3E	6.3E	6.3	6.24
	Expression Signal	1.67	0.65	4.21	4.21	0.62	6.43	0.64	2.91	3.42	0.77	0.55	1.73	0.59	0,59	147		1.47	2.7	2.47	257	2.38							2.81
	ORF SEQ ID NO:	30850	31509	31763	31764	32065	32677	33119			34868		35457	L		27820	1	37630		31065	27401		34786		35026				5 29590
	Exan SEQ ID NO:	18465	18587	18802	18802	19080	18634	20041	20928	21255	١.	L.	1_	L	L	1	200227	24304	1	l	77770	1		1_	1_	1_	L	۱ ۱	16965
	Probe SEQ ID NO:	60	200	8022	6022	8308	6719	7360	8234	8563	9025	9486	9817	10181	10161		20/1	11709	12/4/	12188	,	1/48	ORCO	74.45	919	9913	10171	10615	4224

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Single Exon Probes Expressed in Brain	Top Hit Descriptor No. Source	Rattus norvegicus differentation-associated Na-dependent inorganic phosphate cotransporter (DNPI) mRNA,	TOGGGGIAIG	DATES! TA	1 2	2 4	NT ENT DIMAN	7808 NT	LN OSS	Į.	: !	LV LV		1 EST HUMAN	N.	I N	N DYDYDC+	TOT THEFT	FST HIMAN	T	L		K	EST_HUMAN	EST_HUMAN	Ŋ	EST HUMAN	Ż	EST HUMAN	
Single Exon Probes Expressed in Brain			TOGGSSI	DE LOS			T CI INAMA	NAME OF THE PERSON OF THE PERS					7	HOMAN				THE PARK	Т	Т				Т	T_HUMAN		T		HUMAN	1
	Most Similar (Top) Hit Top Hit Acession BLAST E No.	6.2E-02 AF271235.1	8.2E-02 062191	8.2E-02/D49530 1	6.2E-02 [141453 1	8 2E 02 Bas 4 04 4	6.2E-02 AA778450 1	6.2E-02	6.2E-02 AF217490 1	6.2E-02 AJ242735.1	6 2E.02 AE200350 4	6.2E-02 AE000750.1		8 45 00 040 124 4	8 4E-02 172225 4	8 1E-02	Xooosa 1	6.1F-02 BF974853.4	6.1E-02 BE971853.1	6.1E-02 BE179543.1			6.1E-02 AL163207.2		6.0E-02/AA188730.1	6.0E-02/AE001777.1	6.0E-02/AW968848.1	6.0E-02 AB031289.1	6.0E-02 AA188730.1	
	Expression Signal	1.02	6.31	0.65	1.03	90	0.52	1.65	1.56	1.53	1.74	13.39		2.5 F R0	2.28	7	3.75	0.57	0.57	4.91	1.27	2.27	5.61	0.76	0.76	Ď,	B0:	1.62	6.0	١
	ORF SEQ ID NO:			32656	33292		35092	35214	37039	37316	37781		34037	25697			33986	34388	34389	36560	37787			25559	23060	20002	00107		25559	00000
	Exon SEQ ID NO:		17277	19615	ı			22042	23786	24011	24449	25405	24782	13058	18721	18803	20855	21251	21251	23323	24448	25323	24933	12822	77088	4504	5	15480	12922	10000
	Probe SEQ ID NO:	4304	4542	8699	7527	8846	9243	9380	11095	11320	11865	11989	12394	249	3972	6023	8161	8559	8559	10630	11862	11945	2003	20 20		i cago			2937	

Page 135 of 536 Table 4 Single Exon Probes Expressed in Brain

Top Hit Descriptor	EST84266 Colon adenocarcinoma IV Homo sapiens cDNA 5' end similar to tissue-specific protein	EST84288 Colon adenocarcinoma IV Homo sepiens cDNA 5' end similar to tissue-specific protein	ANABERTANN NIH MGC 68 Homo saplens cDNA clone IMAGE:3876060 3'	Control control to the control of the complete cds	Pos BT055401199-013-b04 BT0253 Homo sapiens cDNA	MASSESSION NEI T GRC S1 Homo sapiens cDNA clone IMAGE:2358873 3' sImilar to contains	When we have a second to the s	Homo sapiens stimulated trans-acting factor (30 KJ/3) (31 AF30) IIINNA	Homo sapiens sumulated transpound track (Control of Control of Con	501815Z74FZ NIT MICC. 30 Halid September CDNA clone IMAGE:1754199 3'	grobbus.X1 Joanes ussus jain i num opmilete genome	Recindmentals americana militariori, conferes genericana de la cione IMAGE:2237362 3'	ts78a06.X1 NCI CCAP GCO nome septems con clone IMAGE:2237362.3	ts/8a06.X1 NCI CCAR GCO name septem of the chain variable region, exons 1-2	Acipenser partial IGLY garle to illustrate and in the chain variable region, exons 1-2	Acipenser been partial IOLV gaile for illiniaring social significant plant in the protein 1, 60 kDe-	EST 180654 JURGIT 1-Cells V Tiding septients College C	EST180654 Jurkat T-cells V Homo sapiens cDNA 5' end similar to similar to heat snock protein 1, oo kDar	fixe	2087-08.1 Stratagers lung calculours 2012 10 Colon 2012 10	Homo sapiens DNA-dependent protein funds e catalytic succinit modernis Processes 3's similar to TR:060298	WRG9h03.X1 Scares_NFL_1_GBC_S1 name squens con your miles and statement of the statement of	RC1-DT0001-290100-012-910 D10001 Homo sapiens cuiva	Mus musculus p53 tumor suppressor gene, exon 10 and 11, parael cus, alta nativaly sprived	Saccharomyces cerevisiae protein tyrosine phosphatase (P.1.7.) gene, comproved as TR-065386	wv34e02.x1 NC_CGAP_OV18 Homo septens cUNA done livraGE.x25143U.5 stitute at 11.00000000000000000000000000000000000	wy34e02.x1 NCI_CGAP_Ov18 Homo saplens cDNA clone IMAGE::2631450 3 similar to 1rt. Occoso	O65386 F12F1.20 PROTEIN ;	R01877609F1 NIH MGC 55 Homo saplens cDNA clone IMAGE:4105994 5'	
 Top Hit Database Source	EST HIMAN	Т	т	ESI HUMAIN	┰	ESI_HUMAN	EST_HUMAN L			\neg	HOMAN		П	T HUMAN		TN	EST HUMAN	Т	EST_HUMAN	T HUMAN		EST_HUMAN		Ν	NT	EST HUMAN		T HUMAN	H LIMAN	בייוטויים ביים
Top Hit Acessian No.	T	Ī		١	1	6.0E-02 AW370211.1	6.0E-02 AI807537.1	74698	74698	_		466495				02 AJ245365.1	2 4 4 3 1 9 7 4	10000	02 AA309797.1	AA128386.1	-02 11431702 NT	AI809273.1	5.9F-02 AW934719.1	02 AF190269.1	-02 AF006304.1	no AWN28748 1		-02 AW028748.1	N 8425508	-02 BF242/48.1
Most Similar (Top) Hit BLAST E Value	100	9.0E-02.A	6.0E-02	6.0E-02 B	6.0E-02 A	6.0E-02 A	6.0E-02	6.0E-02	6.0E-02	6.0E-02	6.0E-02 /	6.0E-02	6.0E-02	6.0E-02	6.0E-02	6.0E-02	20 30 8	0.0E-02	6.0E-	6.0E-02	6.0E-02	6.0F-02	5.9F-02	5.9E-02	5.9E-02	r.		5.9Ε	5.9E	5.9E
Expression Signal		1.48	1.48	0.72	0.69	0.94	22.0	3.07	3.07	233	2.13	0.54	1.17	1.17	1.66	1.66	1	P	0.5	1.69	2.19	2 94	3.87			0.73				0.8
ORF SEQ ID NO:		28639	28640		30370		34868	30524	30525	32842	33355		35007					32659	35660	ŀ	31064	<u>L</u>	25674				3045/	30458	7 34350	
Exon SEQ ID NO:		15986	15986	16378	L	18117		1_	17987		L	L	L	L	L	1	1	22456	22456	ļ		1	1	13035	ł	1	17841	17841		1_1
Probe SEQ ID NO:		3223	3223	3625	5037	5313	200	2710	889	7088	7580	8321	9172	9472	9308	8308		9805	9805	4430B	42487		1230	3 5		2	5123	5123	8515	9351

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Single Exon Done		Γ		_	7	_	_	_						_		_														•		
Single Exon Prosession Nost Similar No. Signal Signal No. Signal Sig	bes Expressed in Brain	Top Hit Descriptor		Mus musculus follistatir-like (Fstt), mRNA	Home saplens nineln (LOC51199), mRNA	Gallus gallus HKC9 telomere junction	Thiobacillus ferrooxidans merC, merA genes and URF-1	Populus trichocarpa CCoAOMT1 gene, expn 1 to expn 5	Wx24c02.x1 NC CGAP Kid11 Homo seplens cDNA clone IMAGE:2544578 3	Wx24oU2.X1 NCI_CGAP_KId11 Homo sapiens cDNA clone IMAGE:2544578 3' qh6801.X1 Socres felat Iliver spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1848687 3' elmiler to	qh86f01xf Scares_fetal_liver_spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE-17448997-2' elmilor to	ga:m13142 COAGULATION FACTOR XI PRECURSOR (HUMAN);	Gallus gallus tyrosine kinase JAK1 (JAK1) mRNA, complete cds	Humen polymorphic microsatellite DNA	Human polymorphic microsatellite DNA	Homo saplens chromosome 21 segment HS21C083	Drosophila melanogaster male fruitless type-A (fru) mRNA, complete cds	no75e11.s1 NCI_CGAP_AA1 Homo saplens cDNA clone IMAGE:1112884 3'	odosbub.s1 NCI_CGAP_Br2 Hamo sapiens cDNA clone IMAGE:1632465 3' similar to WP:C37A2.2	Jonn canions donoming to the second	Chironomus trummi trummi globin VIIA.1 (ctt-7A.1), globin 8.1 (ctt-8.1), globin II-beia (ctt-2heta) non-	uncuonal globin XIII (ctt-13RT), globin XII (ctt-12) and globin XI (ctt-11) genes, complete cds	S 137 8865 MAGE resequences, MAGI Homo sapiens cDNA	Sos taurus Nysozyme gene (cow 3), complete cds	complete reads mixthy for fourth component of complement, complete cds	venopus laews mknA for fourth component of complement, complete cds	Nativas norvegicus mRNA for potassium channel, alpha subunit (kv9.2 gene)	nus muscanus ectz oncogene (Ectz), mRNA	n18b09.y1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC cn18b09 random	118h00 of Name of Liver Table 1	Homo sanions chromosome of a contraction of the Homo sapiens cDNA clone NHTBC cn18b09 random	Pig DNA for SPAI-2, complete cds
Expr ORF SEQ Expression Top Hit Acess	igle Exon Pro			Z	Į.	N I	J.N.	NT	FOI HUMAN	EST HIMAN		EG LICIMAN					T	7					NEW DE						T			
Exam	Sin	Top Hit Acession No.	01.001.00	0/96/00	11433356	000440.4	A 1320624 4	AJZZ30Z1.1				T	T	Ī	T	Ī		T		Γ		Τ			T	Ţ.	81260					\prod
Express		Most Similar (Top) Hit BLAST E Value	A OF 02	3.9E-02	5.9E-02	20 JG 2	5.0E-02	3.0E-02	5.8F-02	5.8E-02	5 8F-02	5.8F-02	5.8F-02.N	5 8F-02	5.8F-02/4	S RE OO	2.0E-02	3.05.02	5.7E-02	5.7E-02 A	5.7E-02	6.7E-02 A	5.7E-02 M	5.7E-02 D	5.7E-02/D	6.7E-02.A	5.7E-02				5.7E-02 AI	5.7E-02 D5
BE Ebon ORI DE 23376		Expression Signal	3.2	1 44	5	5 18	98 0	4.9	6,4	4.95	4.95	2,62	2.99	299	0.67	1 79	200		1.36	1.29	0.97	2.45	1.01	69.0	0.69	1.42	0.64	5	4.42	4.42	1.59	7.24
				Ĺ	l.	 		29687	29688	29879	29880	-	33352	33353	34394			-	28463	28478		29173	-	33203	33204	33880	35606	37000	BONS	37091		
Probe SEO ID NO: 10866 110844 111544 11086 110844 111544 111544 11084 11			1	L	1	l	L	ŀ		L _ I	17245	17270	20247	20247	21257	24590	25386		15819	15834	16448	16535	17371	20115	20115	20749	22401	23810		23810	24012	25213
		Probe SEQ ID NO:	10686	10944	11544	912	2864	4322	4322	4510	4510	4535	7578	7578	8565	12084	12373		3053	3088	3694	3783	4637	7438	7438	8055	9750	11143		11143	11321	12285

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od47f12.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1371119 3' similar to contains Alu repetitive element; contains selement L1 repetitive element; COV0-BN0147-230400-214-g07 BN0147 Homo sapiens cDNA CONA COV0-BN0147-230400-214-g07 BN0147 Homo sapiens cDNA clone IMAGE:2559969 3' similar to gb:X06409 RAF wz34f05.x1 NCI_CGAP_Bm53 Homo sapiens cDNA clone IMAGE:2559969 3' similar to gb:X06409 RAF PROTO-ONCOGENE SERINE/THREONINE-PROTEIN KINASE (HUMAN); qd64g11.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:3453279 5' 601067158F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3453279 5' 601067158F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3453279 5' IMAGE:0CAP_AIM Homo sapiens cDNA clone IMAGE:923245 similar to TR:G769859 G769859 IMAGE:923245 similar to TR:G769859 CF69859 Citrobacter freundii DSM 30040 cyclopropane fatty acid synthase (cfa) gene, partial cds, dlhydroxyacetone Lycopersicon esculentum LE-ACS6 mRNA for 1-aminocyclopropane-1-carboxylate synthase, complets cds zs45c01.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:700416 3' x02c10.x1 NCI_CGAP_Ut2 Homo saplens cDNA clone IMAGE:2656050 3' similar to TR:094979 094979 Hydrocotyle rotundifolia ribosomal protein L16 (rpl16) gene, intron; chloroplast gene for chloroplast product kinase (dhaK), glycerol dehydrogenase (dhaD), transcriptional activator (dhaR), 1,3-propanedio Homo sapiens sodium-dependent vitamin C transporter 1 (SVCT1) mRNA, complete cds Homo sapiens sodium-dependent vitamin C transporter 1 (SVCT1) mRNA, complete cds Homo sapiens elF4E-transporter (4E-T), mRNA Homo sapiens TESTIN 2 and TESTIN 3 genes, complete cds, alternatively spliced Homo saplens fragile 16D oxido reductase (FOR) gene, exons 8, 9, and partial cds Mus musculus second IL11 receptor alpha chain (IL11Ra2) gene, exons 1 and 2 Top Hit Descriptar H.sepiens gene encoding La autoantigen Mus musculus SH3 domain protein 1B (Sh3d1B), mRNA dehydrogenase (dhaT), glycerol dehydratase (dhaB),> Galiid herpesvirus mRNA fragment TROPOMYOSIN ALPHA CHAIN, NON MUSCLE TROPOMYOSIN ALPHA CHAIN, NON MUSCLE Homo sapiens elF4E-transporter (4E-T), mRNA Pan troglodytes apolipoprotein-E gene, complete Mus musculus tuftelin 1 (Tuft1), mRNA Single Exon Probes Expressed in Brain EST_HUMAN EST_HUMAN EST_HUMAN SWISSPROT SWISSPROT EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST HUMAN EST HUMAN Top Hit Database Source Ę 10947034 NT 10947034 NT 눌 눋 Ę 6755902 6755501 Top Hit Acession 5.6E-02 AA482864.1 5.6E-02 AF260225.1 5.5E-02 X97869.1 5.5E-02 L41561.1 5.5E-02 Q01174 5.6E-02 A1983738.1 5.6E-02 A1183583.1 5.6E-02 BE542663.1 6.6E-02 BE542663.1 BE542663.1 BE542663.1 5.6E-02 AW172708.1 AF170911.1 5.6E-02 AA866182.1 5.6E-02 BE008001.1 5.6E-02 AF094455.1 5.6E-02|AB013100.1 AA290599.1 U09771.1 5.7E-02 AF217490.1 5.7E-02 AF261280.1 ġ 5.5E-02|Q01174 5.5E-02 5.5E-02 5.6E-02 (Top) Hit BLAST E Most Similar Value 7.26 3.93 3.05 0.63 1.09 0.66 2.88 3.05 1.12 6.57 3.18 1.57 2.61 Expression 36871 29561 31281 31281 33845 33846 35392 33502 34539 34540 32563 32819 35561 30014 32334 29957 26951 ORF SEQ ÖΝΩ 23622 15370 22363 16932 18370 18370 19861 15972 19754 20388 14265 17330 19535 19742 25283 25387 17382 19327 2071 SEQ ID ÿ 10943 8019 2660 3209 4191 5573 11556 9555 7063 5935 7277 8019 8701 8701 9712 1518 4595 4648 6562 6791 7051 12650 SEQ ID ë

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ngle Exon Probes Expressed in Brain	Top Hit Top Hit Descriptor Top Hit Descriptor			Т	HOMAN		NT Bacillus subtilis complete genome (section 13 of 21); from 2395281 to 2813730	HUMAN		Neurospora crassa ubiquinol-cytochrome c oxidoreductase subunit VIII (QCR8) mRNA, complete ods	7	Т	EST HUMAN IOVO-ST021-0-72-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-	T	T			T Descriptions with the Complete Cds	MT Muse museully a purder tigo gene		NT Helicobacter pulor 26685 service in 134 of the complete genome				Т						
Single Exon Probes Expressed in Brain	Top Hit Acession Defabese Source	1142/1332 NIT		N. C. T. SAAN	ES HUMAN	Z !	<u> </u>	EST HUMAN		NI TOT	ENT LINANI	EST HIMAN	EST HUMAN	EST HIMON		LN L	L	N	 	L	Ę	9695413 NT Lymphocyatis disease vigils 1 com	LN	ΝΤ	SWISSPROT	Т	۲	Į.	Ł	IN TO SE SE TO ANGRE CHARLES	
	Most Similar (Top) Hit BLASTE	5.5E-02	5 4E-03 A 197746	5.4E-02 BE073468 1	R AE 02 105000	5.4E 02 700446.4	5.4E-02.299116.	5.4E-02 AU120889.1	5 45 02 1120700 1	5 4F-02 RE37128	5.4E-02 BF37128	5.3E-02 AW391248.1	5.3E-02 AW391248.1	5.3E-02 T94789 1	5.3E-02 AJ276408 1	5.3E-02 M58417.1	5.3E-02 M58417.1	5.3E-02 AJ276408.1	5.3E-02 M80463.1	5.3E-02 AE000527.1	5.3E-02 AE000527.1	22	5.3E-02 U32832.1	5.3E-02 S78221.1	5.3E-02 P38742	5.3E-02 U10098.1	5.3E-02 X03127.1	5.3E-02 AB022605.1	5.3E-02 AB022605.1	6.3E-02 Y07907.1	
	Expression Signal	1.49	0.094	5.78	0.78	88.0	0.55	1.62	2.03	1.32	1.32	1.28	1.28	14.72	2.47	0.95	0.95	5.51	6.34	1.98	1.98	5.01	F	2.06	0.65	0.7	1.56	0.62	0.62	0.63	į
	ORF SEQ ID NO:	30604			29281		34809	36467	36530	37076	37077	26450	26451	26929	27961	28360	28361	28558	30361	30670	30671	32556	32733		33514		34867	35897	35898		
	SEQ ID NO:		15785	L	16841	L	ŀ	23234	23292	1	23800	13791	13791	14242	15218	15709	15709	15913	17749	18042	18042	19529	19685	18944	8338 8038 8038 8038	86607	21/13	08077	08077	22804	
	Probe SEQ ID NO:	12797	3019	3416	88	8024	8969	10537	10598	11132	11132	1031	<u> </u>	1495	2501	2943	8	32	8	5236	8	8	282	3		3 3	3 3	3 8	75	10156	

Page 139 of 536 Table 4 Single Exon Probes Expressed in Brain

					Sino	III EXOII FION	Single Exon Probes Expressed in State
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Тф Hit Descriptor
	1		6,4	A 3E.00	AF276815.1	Ę	Branchiostoma floridae homecdomain-containing protein Hox13 (Hox13) gene, exon 2 and partial cds
12776		30964	9		31908		Homo sapiens meprin A, alpha (PABA peptide hydrodase) (McF1A) minus
2283	13000				AJ277661.1		Homo sapiens partial LMO1 gene for LIM domain only 1 protein exon 1
3112				5.2E-02	AJ277661.1	NT	Homo sapiens partial LMO1 gene for LIM dollrain off; From; Crd1) mRNA, complete cds
2010	1	29310		5.2E-02	5.2E-02 AF236101.1		Arabidopsis trailians butative divaluations of the second
3021	\mathbf{l}_{-}	L	1.19		6671757 NT		Mus musculus cyclinia inacutation of the complete cds
4245	L	29609	3.02		5.2E-02 U07132.1		Human statute of the capital control of the c
5053	1		6.0		5.2E-02 AA297940.1	HUMAN	ESTITION Communication Communication (CDC54) gene, complete cds
5828	i	31548	0.61	5.2E-02	U14731.1	Ę	Saconaromyces care issue occor. Communication of the second of the secon
	١.		96 0		5.2E-02 AI830965.1	EST_HUMAN	WBURGASTING_CONLYNILE
8018							DNA POLYMERASE PROCESSIVITY FACTOR (TOLIMETS SOLITIONS)
7174	19860	32932	3.13		5.2E-02 P36322	SWISSPROT	BINDING GENE 18 PRO I BIN)
8085	1.		2.19		AL163204.2	Į.	True septens on conditions of the Capsid protein, complete cds
9629	L	35472	1.87	5.2E-02	5.2E-02 D10927.1	Į.	Timils mosely vitas ganomic RNA for Capsid protein, complete cds
96290	1_		1.87		D10927.1	IN.	TUTILIP INCREMENTAL OF THE PROPERTY ASE ALPHA CHAIN
12414	L			5.2E-0	5.2E-02 003030	SWISSPROI	DANE CACE IN ILE DESCRIPTION HIBIT Homo sapiens cDNA clone DKFZp547D073 5'
2364	_			5.1E-0	12 AL 134071.1	ESI HOMAN	Chlamidia trachomatis section 28 of 87 of the complete genome
4179		3 29547	·	6.1E-0	2 AE001301.1	Z	Homo sablens chromosome 21 segment HS21 CO46
496	17685	5	49.38	5.1E-0	12 AL 163246.2	- <u>L</u>	HNV-1 patient 96 from Italy protease (pol) gene, complete cds
6575	19339	32350		5.4	12 AFZBUSOS.1	TOT LI SKANN	OVO-UM0051-250800-350-b08 UM0051 Homo septens cDNA
92/9	17929		4.4	5.4E	02 BF-378625.1	FIGURE I	Human hypoxanthine phosphortbosyltransferase (HPRT) gene, complete cds
816				5.1t	02 M20434.1	NIT	Human hypoxanthine phosphoribosyltransferase (HPRT) gene, complete cds
8151	7 20845			5.1	02 M20434.1	L L	Spodoctera littoralis mRNA for 3-dehydroecdysone 3beta-reductase
8245				5.1E-	02 AJ131966.1	TORGEDIA	KERATIN, TYPE I CYTOSKELETAL 14 (CYTOKERATIN 14) (K14) (CK 14)
878	13 21475	5 34622		5.1E	02 P02533	TOGGGGWG	KERATIN TYPE I CYTOSKELETAL 14 (CYTOKERATIN 14) (K14) (CK 14)
8783	1	6 34623	0	5.1E-	02 P02533	SWISSPRO	Candida gibicans protein phosphatase Sed1 hamolog (SSD1) gene, complete cds
9709	1	35556		5.1E-	02 AF012898.1	1000000	ORIGINATE SPECIFIC PROLINE-RICH PROTEIN APG (PROTEIN CEX)
100	32 22730	35945	1.89	5.1	02 P40603	SWISSPROI	Homo saniena ES18 mRNA, partial cds
10733)	36661		5.16.	02 AF083930.1	Z	Homo sapiens ES18 mRNA, partial cds
10733	<u>.</u>	36662	,	5.1	02 AF083930.1	\ \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Campyopacter leiun NOTC11168 complete genome; segment 3/6
1162	1_	7 37540		5.1E	02 AL 139076.2	Z Z	Chrimis melo polygalacturonase precursor (MPG3) mRNA, complete cds
12421	24797		2.56	5.1E	-02 AF062467.1	Ž	
-							

Page 140 of 536 Table 4 Sindle Exon Pmbas Expressed

Page Except Page Except Page Except Page P			-	Т	7	7	12	_	_	_	_	-	т	_	_	_	_	_			_		_												_	
SEQ ID ID NO: Signal BLASS ID Hit Acession Top Hit Acession	bes Expressed in Brain			Inj73f02.s1 NCL CGAP_Pr10 Homo sapiens cDNA clone IMAGE:998139	Mus musculus fatty acid amide hydrolase gene, excn 10	Bacilius subtilis complete genome (section 1 of 21): from 1 to 213080	SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1/PRP-3) (PRP-2PRP	Optoblem 2012 (PROTEIN C) (CONTAINS: PEPTIDE P.C.)	Min. with the state of the stat	Muss Illustratura Onc-31 like Kinase 2 (C. elegans) (UIK2), mRNA	naturophilus influenzae Kd section 97 of 163 of the complete genome	CASEIN MARKET IT PERSONAL STORY AND AND AND AND AND AND AND AND AND AND	Gallis gallin basin 1987 (CK II)	M	Mich Docen A A Control of the Contro	MECHOLILAMENT I RIPLET L PROTEIN (NEUROFILAMENT LIGHT POLYPEPTIDE) (NFL)	winds musculus res-interacting serine/threonine kinase 3 (Fist3) mRNA, complete cds	90 1044/33F1 NIH_MGC_55 Home sapiens cDNA clone IMAGE:4070101 5	Methanococcus jannaschii section 142 of 150 of the complete genome	INC-CIN-TRANSIENT A PROTEIN	Chicken 28-KDa Wamin D-dependent calcium-binding protein (CaBP-28) mRNA, complete cds	nomo sapiens ABCA1 (ABCA1) gene, complete cds	The County of Anthream States and	24.00 Trilly-1 (JEINTALIOROBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN) 2248812.81 Stratagene hNT neuron (#937233) Homo saplens cDNA clone (MAGE:632928 3; similar to	From the selement contains element MSR1 repetitive element;	A PAGE 72 1 Sources, Testis, NHT Homo saplens cDNA clone IMAGE:728428 3'	Econolist Society and Homo saplens cDNA clone IMAGE:728428 3	Bodg IU.XI IVCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:2632336 3'	BOOGTO.X1 NCI_CGAP_Ut4 Homo sapiens cDNA done IMAGE:2632386 3/	tat etastase II gene, exon 6	rat elastase il gene, exon 6	Archaeoglobus fulgidus section 127 of 172 of the complete genome	memydia muridarum, section 40 of 85 of the complete genome	rackopsis traliana DNA chromosome 4, config fragment No. 59	RANSCRIPTION FACTOR E3	omo saplens chromosome 21 segment HS21C018
Decoration Care Series Capper Similar Cap Hit Aces Capper Signal Cap Hit Aces Cap	gle Exon Pro	Top Hit Detabase Source		EST_HUMAN	ĮN.	L	SWISSPROT	TN	Į	TW	L	TOGGEN		5	Τ	DN-ISS	E INVAN	NAMO -	Τ.	DALL			Т	T.	┰		Т	Т	Т					Т	7	7
Exam	Sin	Top Hit Acession No.	A A 504404 4	AA334104.1	Aruse004.1	289104.1	P02810	Ī	7305610		T			T	T	T		Ī		\	T	T				ı	T	T	T		T	T	T	1		
Exam ORF SEQ Express Sign. 12 24968			A 4E 00	3. 1E-02	3.0E-02	3.0E-02	5.0E-02	5.0E-02	5.0E-02	5.0E-02	5.0E-02	5.0E-02	5.0E-02/	5.0E-02	5.0E-02	5.0E-02	5.0E-02 F	5 0F-02	5 OF -02	4 9F-02 IN	4.9E-02	4.9E-02	4.9E-02 P	4 9E-02 A	4.9E-02 A	4.9E-02.A	4.9E-02 A	4.9E-02 A)	4.9E-02 C	4.9E-02 0	4 OF 021 AT	4 OF 02 A	4 9E-02 A	4 9F-02 P4	4.9E-02 AI	
Example Example No.: No.: No.: No.: No.: No.: 13256 13254 13731		Expression Signal	1 44	1 84	7 9	5.0	3.91	1.28	1.42	1.04	5.83	66.0	0.95	1.3	12.48	1.28	0.45	2.5	3.5	24.03	2.66	2.66	2.53	0.69	0.99	0.99	1.91	1,91	1.9	1.9	140	80	0.77	0.48	0.46	
2012 21 21 21 21 21 21 21 21 21 21 21 21 2		ORF SEQ ID NO:		1				26397			29066		31780		33202	35963	-	37383	-	-	25800	25801	28692		28978	28979	30141	30142	30751	30752	32793	-	34489	36054	36366	
Probe SEQ III NO: I 12677 12677 1483 14666 14678 1473 14770 1677			L	L		L							ı		ш		23167	24074	25246	13028	13158	13158	16043	16311	16334	16334	17519	17519	18091	18091	19733	21205	21344	22839	23140	
		Probe SEQ ID NO:	12678	470	1182		1983	2821	3332	3682	3672	4770	6039	6216	7437	10100	10521	11473	11956	217	380	380	3282	3556	3579	3579	4788	4788	5286	5286	7042	8513	8652	10191	10494	

Page 141 of 536 Table 4 Single Exon Probes Expressed in Brain

					SIID	THE EXOLI PION	Single Exoli Flobes Expressed in State
Probe SEQ ID	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Strnilar (Top) Hit BLAST E	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
:							Homo saniens praprio placental TGF-beta gene, complete cds
11378	23985	37285		4.9E-02	4.9E-02 AF008303.1		Home series CS hox containing WD protein (LOC56884), mRNA
12345	1_		1.77	4.9E-02	8923880 N		
	<u> </u>			00.00	N10364 1	L	Human gamma-B-crystallin (gamma 1-2) and gamma-C-crystallin (gamma 2-1) genes, complete cds
12598				4.95.02	D46474 4	LN	Human mRNA, Xq terminal portion
321				4.8E-02	2 010471.1	H-Z	Human mRNA, Xq terminal portion
322	13123			4.8E-02	02 D164/1.1	F14	Arabidoosis thallana AP2 domain containing protein RAP2.7 mRNA, parttal cds
476	13262		9.96	4.8E-02	02 AF-003100.1	Z	Adamo at Soares senescent fibroblasts NbHSF Homo saplens cDNA clone IMAGE:325611 3 similar to
2027	14007	27735	1.82	4.8E-	02 W51983.1	EST_HUMAN	gb:M30838 LUPUS KU AUTOANTICEN PROTEIN P86 (HUMAN);
7	L			4.8E-	02 X17144.1	Ę	letrahymena rostrata instante non and instante
ğ				4 87	02 754280.1	Z L	S.scrofa gene for skeletal muscle ryanodner terepro
4623				2	14803434 NT	LN	Homo sapiens DKFZP434D222 protein (KEN I.2.), mKNA
5144	4 17863			4.8E-02			Home raniens DKFZP434D222 protein (RENT2), mRNA
5144	L	3 30479		4.8E	11693131 NI	12	MP2 ST0120-231089-012-b02 ST0129 Homo sepiens cDNA
ğ	L		1.32	4.8E	.02 AW388497.1	EST_HUMAN	This are the second and the second a
200	L	1		4.8E	02 AJ001398.1	N	Fugu rupipes (park gene
902	1			A RF	-02 A.1001398.1	LN	Fugu rubripes rps24 gene
905	71 21717	348/1					yz9709.r1 Soares melanocyte 2NbHM Homo sapiens CDNA cione invasication of communications and programment of the communication of the co
. 6	10565	32507	3.83	4.7E	-02 W01153.1	EST_HUMAN	repetitive element
0/31	┸			4.7E	-02 M62752.1	NT	Rat statin-related protein (s.f.) gene, compress con
200	L.		3 8.24		-02 X15543.1	<u>L</u> N	B. Burus miny of the state of t
014	\perp	2 9			4 7F-02 X89211.1	Ł	H. sapiens DNA 1of endogenous renoviral moderns
8852	52 21343				4 7E-02 AR026678.1	F	Gallus gallus Wpkci-8 gene, comptete cas
8875	_1				4 7E 00 X45543 1	Į.	B.taurus mRNA for RF-36-DNA-binding protein
912					DE005007 4	EST HIMAN	601892692F1 NIH MGC_17 Homo sapiens cDNA cighe INACE:4130414
9547	47 22200	35382			4./E-02 BF-303231.1	NAMI IL TOE	we79c10 x1 Soares NFL T GBC_S1 Hano sapiens cDNA clone IMAGE: 234/314 3
9635	35 22287		٥		A1873042	FAI TOWN	Mus musculus ligand of numb-protein X (Lnx), mRNA
10654	54 23345	45 36582			2 6/34303 IVI	IN C	Bos terms natived box protein (bax-6) gene, partial cds
11545	١.	45 37453	1.39		4.7E-02 U73621.1	Z.	Doe towns naired how profein (pax-6) gene, partial cds
11545	L	45 37454		4.7E	4.7E-02 U73621.1	Z	DATA TATAS STATES AND STATES AND SERVICE SERVICES CONA
1	1		0.83	4.6E	-02 BE153583.1	EST HUMAN	Franchis coll K12 MG1655 section 335 of 400 of the complete genome
	L		19 2.91	4.6E	-02 AE000445.1	Z	Estate of Charleton frontal Cortax Homo sapiens cDNA clone IMAGE:1538979 3' similar to TR:P90533
					4 OF A2 A1044255 1	EST HUMAN	amoudu.s.s. Johnson in one comment of the contract of the contains element LTR1 repetitive element;
12	_1	١			4.6E-02 AV727059.1	EST HUMAN	AV727059 HTC Homo saplens dDNA clone HTCBWC01 5'
13	1338 14086	86 26762	3.47	╛	2 UN 12 V V V		

Page 142 of 536 Table 4 Single Exon Probes Fynnessed

SEC ID SEC ID DN SEC Expression Top HI Top HI A Accessed in Driebles Expressed in Driebles Top HI Debtops T				z	٦	7	7			_		_	_	_	_	_	_	_	_	_		Ţ			_	_	_ `	-	. .	-		
December Comparison Compa	bes Expressed in Brain	Top Hit Descriptor	xn24f03.x1 NCI_CGAP_Kid11 Homo saniens cDNA_class_cos_cos_cos_cos_cos_cos_cos_cos_cos_c	Q12849 G-RICH SEQUENCE FACTOR-1:	PMO-HT0339-251199-003-905 HT0339 Homo sapiens cDNA	PM0-HT0339-261199-003-905 HT0339 Homo seplens cDNA	PMO-HT0339-251199-003-905 HT0339 Homo septens cDNA	Mus musculus nucleolar RNA helicase II/Gu (ddx21) gene, complete cds zm92c10.s1 Stretagene ovarian cancer (#897z19) Homo seplens cDNA clane IAAA CE. Exercas and	gb:X03212 KERATIN, TYPE II CYTOSKELETAL 7 (HUMAN); Haplochronis burtoni gonadotropin-raleasina harmana mai O-Ru	gene, complete cds	C.reinhardtil atp2 (аtpВ) mRNA	C.reinhardtil аф2 (афВ) mRNA	9000000x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo seplens cDNA nome IMACE 1720022 A.	similar to contains L1.t3 L1 repetitive element;	Trin-Tri v 0339-060400-009-G12 HT0339 Homo saplens cDNA	out now strategy NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:1624737 31	4V 7128/1 DCA Homo sapiens cDNA clone DCAAZF07 5'	Tuman germline immunoglobulin lambda light chain gene	AETINOIC ACID RECEPTOR BETA (RAR-BETA)	warourg wrus strain M/S. Africal Johannesburg/1976/Ozolin VP35 gene, complete cds	Aarburg virus strain M/S. Africal Johannesburg/1976/Ozolin VP35 gene, complete cds	CALLOCTIE NUCLEAR FACTOR 3-BETA (HNF-3B)	Wella rastidiosa, section 110 of 229 of the complete ganome	onto sapiens chomosome 21 segment HS21C078 Tomo sapiens ASCL3 gene, CEGP1 gene, C11-414 asso, C11-417	ane	omo saplens chromosome 21 segment HS21CDBn	lethanosarcina frisia carbon monoxide dehydrogenase large subunit (cdh.A) gene, carbon monoxide	anydrogenase small subunit (odhlB) gene, complete cds ethanosardna frisia carbon manodde doction	thydrogenese small subunit (cdhiR) dene commissional	abidopsis thallana CCAAT-box binding factor HAP3 homolog	3128167 Cerebellum II Homo sapiens cDNA 5, and claufes to complete cds	A.europaeum mRNA for legitimin-like protein
Exon ORF SEQ Expression Top) Hit Top Hit Acess No. Signal BLAST E No. No. Signal Signal Signal No. Signal Signal No. Signal No. Signal No. Signal No. Signal Signal No. Signal No. Signal Signal No. Signal No. Signal Signal No. Signal Signal Signal No. Signal Signal No. Signal gle Exon Pro			EST HUMAN	EST HUMAN				HOMAN					7	INVAN.	7	TOMAN	7	Т		TOGGGGT	2											
Exon ORF SEQ Expression Top)	EN	Top Hit Acession No.		T		I	T			2.1					T		Ţ		T	T	T		T		1	T					-	
Exon NO: DRF SEQ Express Signa NO: DNO: Signa Signa NO: DNO: DNO: Signa NO: DNO: DNO: Signa NO: DNO: DNO: DNO: DNO: DNO: DNO: DNO:		Most Similar (Top) Hit BLAST E Value	10,	4.0E-UZ	4 RE 02	4 RE 02	4 6F-02	4 6F-02		4.0E-02 A	4.0E-02 X	4.6E-02.X	4.6E-02.A	4.6E-02 B	4.6E-02 A	4.6E-02.A	4 6F-02 X	4.5E-02.P	4.5E-02 A	4 5E-02 A	4.5E-02 P	4.5E-02 AF	4.5E-02 AL		4.3E-02 A)	4.0E-02 AL			4.6E-02 L26	4.5E-02 AF	4.5E-02 AA	4.0E-UZ XB
Exan OR 10 SEQ 1D OR 10 SEQ 1D OR 10 SEQ 1D OR 12 15209		Expression Signal	200	0,7	0 74	E.	1.35	0.89	-	10.	0.01	10.0	1.47	269	49.4	1.88	3,98	1.72	1.11	1.11	4.57	3.76	3.66	20 4	12.0		0.61	3	19.0	OR'	4.07	V.*01
8 0 :: 8 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1				L							31884		32659	34386	37286			25866	26612	26813	27247	27568	29102	31885	32182		32550	27854	3440E	35600	35860	10000
8 0 : 8 7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				Ľ	L	L		i			1	1	19617	21246	23986	24744	24985	13220	13948	13948	14537	14834	16463	18915	19183		19523	19523	2008	22499	22648	
		Probe SEQ IC NO:	2492	2811	3325	3487	4103	6121	5647	6136	6136		6702	8554	11379	12325	12705	\$\$	1196	1198	1797	2103	3710	6137	6416		8778	67.79	8282	9849	10000	-

Page 143 of 536 Table 4 Sindle Exon Probes Expressed in Brain

Single Exon Probes Expressed in Brain	Top Hit Descriptor	Gallus gallus mRNA for alpha1 integrin, complete cds	Homo sapiens ret finger protein-like 3 (RFPL3), mRNA Homo seriens cDNA clone IMAGE:632493 5'	13f11,r1 Stratagene hN I neuron (#50/1255) I tonic of the properties of the properti	601652154F1 NIH_MGC_82 Homo sapiens cDNA cigne IMAOE3930000	HYPOTHETICAL PROTEIN (ORF 2280)	QV2-PT0012-010300-070-902 PT0012 Homo sapiens cDNA	Myxococcus xanthus serine/threonine kinase Pkn10 (pkn10) gene, camplete cas	Homo sapiens S164 gene, partial cds, PS1 and hypothetical protein ganas, complete cus, and complete cu	partial cds.	partial cds	Ows ares COART of mainteen and mainteen are 9 (MMP-9) mRNA, partial cds	Carils latinification matrix metalliproteinase 9 (MMP-9) mRNA, partial cds	Central Harring Industry Months State of State o	Hepatitis E virus strain HEV-US2 polyprotein (ORF1), (ORF3), and capsid protein (ORF2) genes, complete	cds e-23104 r1 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:897631 5'	Homo saniens mRNA for KIAA1493 protein, partial cds	601878746F1 NIH MGC_55 Homo sapiens cDNA clone IMAGE:4107418 5	Morone saxatilis myosin heavy chain FM3A (FM3A) mRNA, complete cds	AV704878 ADB Homo sapiens cDNA clone ADBAOH08 5	Homo sapiens chromosome 21 segment HS21C010	Homo sapiens promyelocytic leukemia zinc finger protein (PLZF) gene, complete cus	Grapevine fanleaf virus coat protein gene, partial cds	PLECTIN	PLECTIN	ns69c12.s1 NCI_CGAP_Prz Homo sapiens cDNA clone IMAGE:1188886	Homo sapiens desmocollin 3 (DSC3) gene, complete cds, alternatively spliced	Hisapiens NCAM mRNA for neural cell adhesion molecule	H.sapiens NCAM mRNA for neural cell adhesion molecule	AU123327 NT2RM2 Homo sapiens cDNA clone NT2RM2000020 o	AU123327 NT2RM2 Homo sapiens cDNA done N 12RM2000020 3	
e Exon Probes	Top Hit Database Source	Š NT		HUMAN	Г	Г	Т	M TZ	Т	ă I			N N	1444411	NEW DE LES	\neg	NICANOL -	MAN	Т	T HIMAN	T			ISSPROT	1.	T				T HUMAN	ı	1
Singl	Top Hit Acession No.	4 EE 02 AB000470 1	8013		4 AE 00 BE 070733 1		١		02 AF 139 100.1	02 AF109907.1	02 AF109907.1		02 AF095824.1	02 AF095824.1	-02 AA736969.1	-02 AF060669.1	-02 AA496739.1	-02 AB040928.1	-02 BF241245.1	-02 AF-003249.1	4.3E-02 AV / 048/8.1	4.3E-UZ AL 100210.2	4.3E-02 AF000300.1	4.3E-02/011706.1	4.3E-02 F30427	4.3E-02 P3042/	4.3E-02 AA652200.1	-02 AF283339.1	4.3E-02 X55322.1	4.3E-02 A33322.1	4.ZE-02.AU 120327.1	100 1000
	Most Similar (Top) Hit BLAST E Value	A EE 02 A	4.5E-02	4 5F 02 A	3 60 37 7	4.45-02	4.4E-02 P31300	4.4E-02/	4.45-02/	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02	4.35-02	4.3E-02	4.3E-02	1					4.3				_
	Expression Signal		200	200	18:0	5.52	5.42	2.4	1.95	1.24	1.24	2.28	0.64	0.64	2.14	2,62	2.78	2.57														2.32
	ORF SEQ ID NO:		35976	2109/	30/33	1		27953	29024	29945	20048		32766			36936	37104	L	Ц		Ц	28830									5 26238	8
	Exon SEQ ID NO:		22764	24045	2520	13025	14821	1	16384	17319	L	L	1_		<u> </u>	23679	L	1_	25408	13536	15287	16180	16404	1	19173	1	١_	1 21104			1	13616
	Probe SEQ ID		10118	12154	12537	213	2089	2494	3631	4584	7627	2037	201	7018	8650	11007	11187	11890	12067	783	2573	3423	365	5152	ğ	6404	9633	8411	8700	8700	803	8

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	Top Hit Descriptor	w34g01.XI NCI_CGAP_PIt1 Homo saplens cDNA clone IMAGE:2545684 3' similar to TR-O63394 O63394	LI RE I NOPOSON, ORFZ MRNA ;contains L1.13 L1 L1 repetitive element ;	enome; segment 4/5			601150933F1 NIH_MGC_19 Hamo saplens cDNA clone IMAGE:3503505 6	ulation (GSG1) gene, complete cds	ulation (GSG1) gene, complete cds	Homo septems cytochrome P450 polypeptide 43 (CYP3A43) genes, partial eds; cytochrome P450 polypeptide 44 (CYP3A44) and cytochrome P450 polypeptide 7 (CYP3A47) genes, complete eds; and cytochrome P450 polypeptide 7 (CYP3A77) genes, complete eds; and cytochrome P450	Homo saplens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide 4 (CYP3A4) and cytochrome P450 polypeptide	polypeptide 5 (CYP3A5) gene, partial ods	the CDNA close MAA CE 2000010 FT	856 (katA) dene complete ode	ALPHA-ACTININ 3, NON MUSCULAR (F-ACTIN GROSS LINKING PROTEIN)	T-BRAIN-1 PROTEIN (T-BOX BRAIN PROTEIN 1) (TBR-1) (TES-56) on33b11.s1 NC_CGAP_Lu6 Homo septions cDNA chine IMA CE 1455444 31 -1 "	URSOR (HUMAN);	Homo sepiens cDNA	Homo sapiens cDNA	inotein gene, complete cds	Mens cDNA clone IMAGE:2510850 3'		the complete genome	Jomo sapiens cDNA	ans cDNA clane IMAGE:3343856 5	ins cDNA clone IMAGE:3343856 5'	iumsic protein 1a	or the complete genome
Single Exon Probes Expressed in Brain	Hit base roe		7	1	Ť	Т	Т	Saccharomyces cerevisiae general sporulation (GSG1) gene, complete cds	Т		Homo saplens cytochrome P450 polypepti		_	П				Т	T	Т	T	Chimile septents n.P.S.I gene, intron 5	T	Ŧ	Т	A.thaliana mRNA for plasma mambrana harmana ha	Ureaplasma urealyticum section 33 of 80 of the	duos eu la colla colla colla company
ngle Exor	Top Hit Database Source	100		SW/Scoop	SWISSEDET	CAT LE	ES L' HUMAN	Z L	EST HIMAN	E			EST_HUMAN	NT	SWISSPROT	SW ISSPROT	EST HUMAN	EST HUMAN		FST HIMAN	a .	LZ	EST HIMAN	EST HIMAN	EST HUMAN			
<u> </u>	Top Hit Acess No.	4.2E-02 AW003645 4	-02 Al 445066 1	-02 P23091	P23091	-02 RE28260E 4	-02 1126674 1	02 1128874 1	02 BF342995.1					52.1	72 P05095		4.2E-02 AAB/6118.1	T		Γ				T	T.		AE002132.1 NT	11 11 00000
	Most Similar (Top) Hit BLAST E Value	4.2E-02	4 2F-02	4.2E-02	4.2E-02	4 2F-02	4.2E-02	4 2E-02	4.2E-02	4.2E-02		4.2E-02 /	4.ZE-0ZI	4.2E-02 /	4 2E 02		4.2E-02.4	4.2E-02.B	4.2E-02.A	4.2E-02 AI983494.1	4.1E-02 A	4.1E-02 A	4.1E-02 A	4.1E-02 BE	4.1E-02 BE	4.1E-02 X7	4.1E-02 AE	1 4 11 00
	Expression Signal	1.35	1.02	101	2.43	0.7	1.83	1.83	2.32	0.68		88.0	3	4.7	1 22		2.02	2.54	1.68	3.43	1.24	1.04	7.52	0.82	0.82	0.67	1.25	200
	ORF SEQ ID NO:	26315		27213	29047	29471	29648	29649	30060	31231		30517	32400	34547	36930	36808	37231	37232	37402		26916	28131		31262	31263		32742	33174
	Exen SEQ ID NO:	13645	14457	14513	16408	16843	17023	17023	17429	18328	0000	17962	20103	21402	22712	23645	23939	23939	24090	25335	13281	16392	17175	18353	18353	19527	16031	06007
	Probe SEQ ID NO:	876	1714	1771	3865	4100	4284	4284	4695	6630	7830	9888	742A	87.10	10064	10969	11278	11278	11489	_					┙	L	Page 1	

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Most Similar (Top) Hit Top Hit Acession Database BLASTE No. Source	Mus musculus provinal retrovinal insertion in the cGMP-phosphodiesterase (rd beta PUE) gene, introri 1, with the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' LTR the provinal insert encompassing the env pseudogene (3' end) and 3' encompassing the env pseudogene (3' end) and 3' encompassing the env pseudogene (3' end) and 3' encompassing the env pseudogene (3' encompass	4 1E-02 AF026198.1	4.1E-02 P34687 SWISSPROT	4.1E-02 AA372398.1 EST_HUMAN	4.1E-02 AJ271909.1	4.0E-02 AB040904.1 NT	127 4.0E-02 L11910.1 NT Human reunoxascums susceptuming special control of the co	TW 17040903 A CO TO 4		0.93 4.0E-02 BF110434.1 EST HUMAN R29124.1.1	6.57 4.0E-02 L23838.1 NT cds		LN	4.0E-02 AB000381.1 NT		4.0E-02 BF679376.1 EST_HUMAN	02 AJ000941.1 NT	02 D43949.1 NT	4.0E-02 AJ001018.1	4.0E-02 AJ001056.1	3.9E-02 BF516149.1	3.9E-02 P41047 SWISSPROI	2.4 3.9E-02 AJ403386.1 N1 Home seniens succinate dehydrogenese complex, subunit C, integral membrane protein, 15kD (SDHC)		3,9E-02	3.9E-02 8924019 NT
Expression Signal	0.08																	1.21						10		
ORF SEQ ID NO:	33265			34899	L		29170	Ì	30759	31866	33366		33440	1	34451		35406	4	6					Ľ	29487	
Exon SEQ ID NO:	20173	i _	20329		1	1	1	1	18100	18898		1	1	20330	1		L	L	L	١_	8 13856	L	14689	15415	1	1
Probe SEQ ID NO:	7502	36	7665	8041	12728	3238	3780		9239	6120	1 29	7650	7666	7666	7120	954	9567	9884	11778	12053	1098	132	1954	2300	4118	4118

Page 146 of 536 Table 4

					is	ngle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2408	18207	30913	. 0.55	1	3 9F_02 DE0608 4	E.A.	
6408	18207		0.55		OS DAGGO 4	ž į	Kat gene tor chotecystokinin type-A receptor (CCKAR), complete cds
5644	ŀ		10,	3.9E-	BE968841 1	ENT LIMANI	Rat gene for cholecystokinin type-A receptor (CCKAR), complete cds
5766	18557		0.95	3.9E-	BF675203 1	EST HIMAN	6004387254 NIII WGC 74 Homo sapiens cDNA clone IMAGE:3833642 57
6957	19439		1.18	3.9E-	02 BE271437 1	EST HIMAN	RM14A720E1 NIT MGC_83 Home sapiens cDNA clone IMAGE:4274910 5'
7739	20435		1.14	3.9E-02	BF239613.1	EST HIMAN	BOTTONESSE INITIALIZED FOR SECTION SECTION OF THE PROPERTY OF
7959	1 1	33778	0.79	3.9E-02	02 AJ229041.1	L	Homo sanjane 050 Uh sanifa hatta A III 4
7959	20654	33779	0.79	3.9E-02	12 A 1229041 1	5	Home carlos of the control will be served AML and CBK1 on chromosome 21q22; segment 1/3
11386	20396	33511	2	3.9E-02	12 P48778	POGGGIVIS	I WITH SALIEUS BUS ND CONTIG Detween AML1 and CBR1 on chromosome 21q22, segment 1/3
11913	L		15.38	3.9E-02	2 AB042553.1	TN TN	Fells cetus C. CSE cons. Co. C. C. C. C. C. C. C. C. C. C. C. C. C.
							ous cause octor gene for granulocyte colony-ethnulating factor, complete cds
					_		Human germline T-cell receptor hela chain TCBBV/4784.44T TCDBV/984
07307							TCRBV19S1P, TCRBV16S1, TCRBV16S1A1T HVR MIC TCRBV28S1P TCRBV28S1P,
25043	24883		1.83	3.9E-02	3.9E-02 U66061.1	L	TCRBV3S1, TCRBV4S1A1T, TRY4, TRY6, TRY7, TRY8, TCRBN TCRB NS 1 TCRB NS 1 TCRB NS 1
12666	25223		- t	i c	000000	!	Mus musculus chromosome X contgB; X-linked lymphocyte regulated 6 gene. Zinc finaer protein 275. Zinc
1845		27304	2.0	3.95-02	3.9E-02 ALU49800.2	Z	tinger protein 92, mmxq28orf
2114	L	+	22.	3.9E-02		ES HOMAN	601610891F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912215 5'
4878	L	90000		2010	0.00-02 0.02019/3.1		Homo sapiens partial steerin-1 gene
F354	L	30840		3.8E-02		EST_HUMAN	AU124122 NT2RM2 Homo saplens cDNA clone NT2RM2001698 6'
200g	L	26772	- 3	3.8E-UZ M11228.1		Į.	Human protein C gene, complete cds
7218	J	32078	1.32	3.8E-02 P10284	00011000	SWISSPROT	HOMEOBOX PROTEIN HOX-B4 (HOX-2.6)
8562	L		133	3 8E.02 Menaze	00/000	Z	Homo sepiens A IP-binding cassette, sub-family A (ABC1), member 8 (ABCA8), mRNA
10549	L	36481	2.62	3.8F-02/4		2 12	Human von Willebrand factor gene, exons 23 through 34
971	13736	26401	4.94	3.7E-02	3.7E-02 P19137	SWISSPROT	LAMININ ALPHA-1 CHAIN BEECHES / AMININ A PARK
-	,			 			Homo Sapiens plasma membrane calchim A TDasa ionform 4 (A TDSD4)
1367	14115	26790	0.91	3.7E-02		TN	partial dis
25.50	4536	27698	3.84	3.7E-02/A		EST_HUMAN	w85e08x1 NGI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:23a4803 2
700	15298	28034	0.92		31.1	7	Homo saplens mRNA for KIAA0718 protein, partial cola
3043	13811	28457	0.9	3.7E-02 P		SWISSPROT	EOMESODERMIN
) 104	15813	28458	2.99		BF312963.1	EST_HUMAN	601896233F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE-4126584 5
3447	16203	-	1.17	3.7E-02	6680541 NT		Mus musculus potassium large conductance pH-sensitive channel, subfamily M, alpha member 3 (Konma3), mRNA
8269	25422		0.83	3.7E-02	AP000063.1		Aeropyrum pernix denomic DNA section 8/7
							The feet of the country of the

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					ino	אום באטוו ר ועג	Single Excit Floors Explassed in Diam
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7592	1_	33368	0.56	3.7E-02	2 AE003975.1	NT	Xylella fastidiosa, section 121 of 229 of the complete genome
9914			-	3.7E-02		EST_HUMAN	al55c09.s1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone 1360912 3'
11964		37811	3.86	3.7E-02	3.7E-02 BF124974.1	EST_HUMAN	601762117F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:4024973 5
12603	25193	30813		3.7E-02	11418392 NT	NT	Homo sapiens solute carrier family 22 (organic cation transporter), member 1 (SLC22A1), mRNA
3646	16399	29039	1.38	3.6E-02	3.6E-02 X73221.1	IN	H.vulgare Ss1 gene for sucrose synthase
3654	16407	29046	0.88	3.6E-02	AL096806.1	IN	Homo saplens genomic region containing hypervariable minisatellites chromosome 10[10q26.3] of Homo saplens
F344	1			3 RE_03		Ŀ	C.glutamicum gap, pgk and tpl genes for glyceraidehyde-3-phosphate, phosphoglycerate kinase and inceenhosphate isomerase
3	5			70.0	Ī		Control of the second of the s
5341	18144	30823	0.58	3.6E-02	2 X59403.1	NT	C.glutamicum gap, pgk and tpi genes tor giyceraldenyde-3-prospinate, prospinoglycerate kinase and triosephosphate isomerase
5413	18212		0.64			IN	Homo saplens RU2AS (RU2) mRNA, complete cds
6607	[32382	24.5			EST_HUMAN	CM2-EN0013-110500-192-b10 EN0013 Homo sepiens cDNA
6607	19370	32383	5.47	3.6E-02		EST_HUMAN	CM2-EN0013-110500-192-b10 EN0013 Homo saplens cDNA
6985	19678		2.5	3.6E-02		NT	Chromatium vinosum sulfur globule protein Cv2 precursor (sgp2) gene, complete cds
7208	19891	32967	2.76	3.6E-02	3.6E-02 AA714521.1	EST HUMAN	nw20e05.s1 NCI_CGAP_GC80 Homo saplens cDNA clone IMAGE:1241024 3' simitar to gb:J00314_ma2 TUBULIN BETA-1 CHAIN (HUMAN);
7533	20203			3.6E-02		EST_HUMAN	MR0-HT0158-030200-003-b08 HT0158 Homo saplens cDNA
9281	21958	35130	1.72	3.6E-02	2 U20608.1	LΝ	Dictyostelium discoldeum unknown spore germination-specific protein-like protein, orf1, orf2 and orf3 genes, complete eds
		L					Dictyostelium discoldeum unknown spare germination-specific protein-like protain, orf1, orf2 and orf3 genes,
9291	21958	35131	1.72	3.6E-02	2 U20608.1	NT	complete cds
9512	L			3.6E-02	2 BF347586.1	EST_HUMAN	602020463F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4156116 5'
11135	23803	37080	1.4	3.6E-02		EST_HUMAN	601820416F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4052570 5'
11135	23803	37081	1.4	3.6E-02		EST_HUMAN	601820416F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4052570 5'
11852	1		1.46	3.6E-02	1		qk48b09.x1 NCI_CGAP_Co8 Homo sapiens cDNA clone IMAGE:1872185 3'
875		26314	1.08	3.5E-02		INT	Drosophila melanogaster tiggrin mRNA, complete cds
888	13751	26413	1.39	3.5E-02		TN	Homo sapiens microsomal epoxide hydrolase (EPHX1) gene, complete cds
1558	14303		1.55	3.5E-02		EST_HUMAN	602085136F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249377 5
1556	14303			3.5E-02		EST_HUMAN	602085136F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249377 5'
4188	16929	29559		3.5E-02	3.5E-02 AE001773.1	NT	Thermotoga maritima section 85 of 136 of the complete genome
4281	17020	29647	1.27	3.5E-02 P53780	P53780	SWISSPROT	CYSTATHIONINE BETA-LYASE PRECURSOR (CBL) (BETA-CYSTATHIONASE) (CYSTEINE LYASE)
6127		31873	1.77	3.5E-02	3.5E-02 J01238.1	NT	Maize actin 1 gene (MAc1), complete cds

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			_	-	_	_	-	-	_		_	_																		
Single Exon Probes Expressed in Brain	Top Hit Descriptor	yp44e05.r1 Soares retina N2b5HR Homo septens cDNA clone IMAGE:190256 5' similar to contains Alu repositive element:	601644701R2 NIH MGC 56 Homo septens cDNA clone IMAGE:3020737 3	Litactis MG1363 grpE and chalk genes	601344661F1 NIH, MGC, 8 Homo sapiens oDNA clone IMAGE:3877634 5'	PM1-CT0326-291299-002-h03 CT0326 Homo sapiens cDNA	PM1-CT0326-291289-002-h03 CT0326 Homo saplans cDNA	601178765F1 NIH MGC 20 Homo seplems cDNA closs IMAGE 3843833 R	Homo sapiens mRNA for FLJ00013 protein, partial eds	Homo sapiens mRNA for FLJ00013 protein, partial cds	Homo sepiens mRNA for FLJ00013 protein, partial cds	Homo saplens mRNA for FL.100013 protein partial ods	xx28d07.xt Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2814253 3' sImiler to SW:C211 HUMAN PSS801 PUTATIVE SI IREACE SI VCAPBOTEIN CAROLES POPULIATIVE SI IREACE SI VCAPBOTE SI VCAP	Homo sablens hyadhetical protein F. 113220 (F. 11322n)	yo20e06.r1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:81250 5' similar to contains	Homo sablens chromosome 21 segment HS21Cn08	PC3-FN0155 DROYOU 011-40 ENDING	RC6-UM0015-210200-021-A10 UM0015 Home senions and	M.musculus S-entitoen cene promoter readon	LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (I A ALITOANTIGEN HOMOLOG)	Caenorhabditis elegans mRNA for DYS-1 protein, partial	Human lysyl oxidase-like protein gene. exon 3	WIS9004.X7 NCI_CGAP_Bm25 Hamo sapiens cDNA clone IMAGE:243303131	nu70f08.s1 NCI_CGAP_Aiv/ Homo sepiens cDNA clone IMAGE:1218071 similar to contains Alu repetitive element; contains element MER25 MER25 repetitive element;	жд04f11.s1 Stratagene muscle 637209 Homo sapiens cDNA clone IMAGE:628749 3' similar to	INCOUNTES CICITY STORY IN THE STATE OF THE S	PPT GPVVISDITEESVTLKWEPPKYDGGSQVTNYILLKRETSTAVWTEVSATVARTAMAKVAMKI	0299h08.x1 Soares_parathyroid_tumor_NbHPA Homo saplens cDNA clone IMAGE:1683519.31	#75e08.s1 Soares_tests_NHT Homo saplens cDNA clone IMAGE:728198.3"	Cricetulus griseus CYP2A17 mRNA for cytochrome P450 2A17, complete cds
igie Exon Pro	Top Hit Database Source	EST HUMAN	EST HUMAN	LZ LZ	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	L	LN	NT	N	EST HUMAN	Į.	NAMI H FSH	L	EST HIMAN	EST HUMAN	LN	SWISSPROT	Z.	Ν	EST_HUMAN	EST_HUMAN			EST_HUMAN	EST_HUMAN	EST_HUMAN	NT
lio	Top Hit Acession No.	-02 H28951.1	-02 BE958970.1	-02 X76642.1	-02 BE561042.1	-02 AW861641.1	-02 AW861641.1	-02 BE276948.1	-02 AK024424.1	3.4E-02 AK024424.1	-02 AK024424.1	02 AK024424.1	02 AW 274020.1	11345459 NT	02 157160.1	2	I							_			1			3.3E-02 AB035867.1
	Most Similar (Top) Hit BLAST E Value	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02 AI869629.1	3.4E-02			3.4E-02/	3.4E-02/	3.3E-02 /	3.3E-02 A
	Expression Signal	0.78	2.7	1.45	0.5	1.82	1.82	5.69	1.14	1.14	6.47	6.47	2.92	7.14	2.06	4.1	0.7	3.19	2.41	3.59	1.2	4.73	3.25	1.36			5.97	0.83	9.61	17.88
	ORF SEQ ID NO:		34357	35762			37389		25973		25973	25974	26448	-	27849	28831	29145	29291	28922		30349	30558		34482					-	26559
	Exon SEQ ID NO:	20572	21213	22566	22813	24078	24078	25234	13346	13346	13346	13346	13789	13936	15112	16181	16509	16650	17294	17723	17740	17923	20853	21338			21506	22330	13161	1388
	Probe SEQ ID NO:	7877	8521	9917	9962	11477	11477	12596	564	28	565	565	1029	1184	2391	3424	3757	3900	4659	2000	5019	8754	8159	8646			8814	9678	88	1143

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Stmiler (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1489	14236	26923	1.16	3.3E-02	3.3E-02 L16870.1	NT	Homo sapiens cytochrome P4502C18 (CYP2C18) gene, exons 2 and 3
1635	14381	27068	1.47	3.3E-02		NT	Homo sapiens skeletal muscle LIM-protein 1 (FHL1) gene, complete cds
1732	14474		1.29	3.3E-02	.1	IN	Aquifex aeolicus section 32 of 109 of the complete genome
2077	14809		2.48		3.3E-02 R09112.1	EST_HUMAN	y/25c09.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:127888 5'
2463	15171		1.31		6755862 NT	NT	Mus musculus tumor rejection antigen gp96 (Tra1), mRNA
4158	14381	27068	2.44		.1	NT	Homo sapiens skeletal muscle LIM-protein 1 (FHL1) gene, complete cds
4435	17171	L	1.78		6755862 NT	TN	Mus musculus tumor rejection antigen gp96 (Tra1), mRNA
6336	19106	L	27.36			EST HUMAN	601853910F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4073787 5'
6336	19106		27.36		BF245995.1	EST_HUMAN	601853910F1 NIH_MGC_57 Homo saplens cDNA clone IMAGE:4073787 5'
7408	20085	33169	0.63	3.3E-02	AF124162.1	NT	Nicottana plumbaginifolia molybdopterin synthase sulphurylase (cnx5) gene, partial cds
9222		35071	0.74		BF115621.1	EST_HUMAN	7m92d04.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:3562423 3'
9222	L	35072	0.74	3.3E-02	2 BF115621.1	EST_HUMAN	7m92d04.x1 NCI_CGAP_Bm23 Homo saplens cDNA clone IMAGE:3562423 3'
8324		35162	0.66	3.3E-02	2 AA488202.1	EST HUMAN	ad08f09.sr Soares_NbHFB Homo saplens cDNA clone IMAGE:877673 3' simitar to gb:X70944_cds1 MYOBLAST CELL SURFACE ANTIGEN 24.1D5 (HUMAN);
	1						ad08f09.s1 Soares_NbHFB Homo sapiens cDNA clone IMAGE:877673 3' similar to gb:X70944_cds1
9324	_		0.68	3.3E-02	2 AA488202.1	EST HUMAN	MYOBLAST CELL SURFACE ANTIGEN 24.1D5 (HUMAN);
11065		37008	3.63	3.3E-0	-	EST HUMAN	602247171F1 NIH_MGC_62 Hamo septens cDNA ctone IMAGE:4332497 5
12142	24630		3.24			EST_HUMAN	ye49f11.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:121101 5'
12259	24704		1.52		.1	NT	Mus musculus EIF4H gene, partial cds; LIMK1 gene, complete cds; and ELN gene, partial cds
12288	24718		2.92			NT	Human interleukin 11 (IL11) gene, complete mRNA
129	12944	25588	0.74	3.2E-02	3.2E-02 AJ002005.1	NT	Oryctolagus cuniculus gene encoding ileal sodium-dependent bile acid transporter
1104	13861	26520	7.01	3.2E-02		NT	Drosophila malanogaster haat shock protein 68 (hsp68) gene, hsp68d allele, complete cds
1104	13861	26521	7.01	3.2E-02	75.1	NT	Drosophila melanogaster heat shock protein 68 (hsp68) gene, hsp68d allele, complete cds
2112	14843		3.01	3.2E-02 P28955		SWISSPROT	LARGE TEGUMENT PROTEIN
3131	15896		10.08	3.2E-02		EST_HUMAN	601442431F1 NIH_MGC_65 Hamo saplens cDNA clone IMAGE:3846727 5'
3701	16454		0.92	3.2E-02	3.2E-02 AL163203.2	NT	Homo sapiens chromosome 21 segment HS21C003
3942	16692		1.64	3.2E-02		ΤN	S.cerevisiae chromosome IV reading frame ORF YDL055c
3942	16692	28331	1.64	3.2E-02		ΝT	S.cerevisiae chromosome IV reading frame ORF YDL055c
4188	16934		14.21	3.2E-02	2 X94768.1	NT	H.saplens RP3 gene (XLRP gene 3)
4716	17448	30081	3.42	3.2E-02	2 AF114182.1	ΤN	Saxifraga nidifica maturase (matk) gene, chloroplast gene encoding chloroplast protein, partial cds
4894 4984	17621		1.09	3.2E-02		LΝ	Mus musculus MHC class III region RD gene, partial cds; Bf, C2, G9A, NG22, G9, HSP70, HSP70, HSC70t, end smRNP genes complete cds; G7A gene, partial cds; and unknown genes
5448	l	31135		3.2E-02	3.2E-02 X68709.1	TN	S.griseocameum whiG-Siv gene

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Single Exon Probes Expressed in Brain	t Top Hit Acession Database Source	12 X68709.1	02 M32437.1 NT Ratipolyomentus left Junction in cell line W 98.14	yd33h12.s1 Soares fetal liver spleen 1NFLS Home septens cDNA clone IMAGE:110087 3' similar to contains 02 T89367.1 EST_HUMAN Alu repetitive element; on the contains and the contains are contained as a contains and the contains are contained as a	2 AF173845.1	11424049 NT	10565 NT	FZ	EST_HUMAN	3.2E-02 Al278971.1 EST_HUMAN qm17b04.x1 NCI_CGAP_Lu5 Homo septens cDNA clone IMAGE:1882063.3	FST HIMAN	3.2E-02 U96762.1 INT Macaca mulatta chemokine receptor CCR5 mRNA, complete cds	3416 NT	SWISSPROT	6671564 NT	2 Z50097.1 NT	Ł	EST_HUMAN	Т	Netsserta meningtidis DNA for region 2 (fhaB- and fhaC-homologs, unknown genes) and flanking genes, strain FAM18	L HUMAN	Į.	3.1E-02 6754241 NT Mus musculus histidine rich calcium binding protein (Hro.), mRNA	3.0E-02 AF187125.1 NT Pityoktelnes minutus cytochrome oxidase I gene, partial cds. miliochondrial gene for mitochondrial broduct	EST_HUMAN		1 EST_HUMAN	EST_HUMAN		NT	12] AB048783.1 NT Homo saplens mRNA for KIAA1573 protein, partial cds
		3.2E-02 X687	3.2E-02 M32437.1	3.2E-02 T893	3.2E-02 AF1	3.2E-02	3.2E-02	3.2E-02 AF10	3.2E-02 AI27	3.2E-02 A127	3 2E-02 AA7	3.2E-02 U967	3.1E-02	3.1E-02 P18845	3.1E-02	3.1E-02 Z500	3.15-02 U78104.1	3.1E-02 AA27	3.1E-02 BF68	3.1E-02 AJ39	3.1E-02 BE96	3.1E-02 AF03	3.1E-02	3.0E-02 AF18	3.0E-02 AA40	3.0E-02 AF24	3.0E-02 AW8	3.0E-02 AA38	3.0E-02 AF28	3.0E-02 AF28	3.0E-02 AB04
	Expression (T	1.83	3.13	33.46		0.64	4.64			1.21	4.07		2.14	_		1.34	1.13	2.12			0.48					2.78			5.83		3.43
	ORF SEQ ID NO:	31138	32196		32279	33436	34030		34978	34979		36114		26702	27332		30206		31268	31338	34677	35778	37689	<u> </u>	28040	28038			30318	30319	
	SEQ ID NO:	18247	19199	19200	19278	20326	20893	21531	21813	21813	22605	22904	13986	14032	14622	14703	17990	18081	18358	25072	21532	22579	24356	14366	15304	16398	16480	16679	17714	17714	18112
	Probe SEQ ID NO:	5448	6431	6432	6513	7662	8199	8839	9125	9125	8967	10256	1237	1282	1885	1967	5182	5276	5561	928	8840	9331	11765	1619	2590	3645	3728	3929	4991	4891	5307

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	Top Hit Descriptor	za89a10.r1 Soares fetal iver spleen 1NFLS Homo sapiens cDNA clone IMAGE:294906 5' similar to contains element TAR1 repetitive element;	za39a10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:294906 5 similar to contains element TAB1 reportitive element:	Contract camp mRNA for inducible nitric oxide synthase (INOS gene)	CONTINUE CADIO III M.C. 71 Home saniens CDNA clone IMAGE:3913848 5	00 10 12200F1 WITH MOO. 71 Home canions cDNA clone IMAGE:3913848 5	United States and less factor of kappa light polyneride gene enhancer in Broells 1 (NFKB1) gene, complete	ווס פקוופווא וותופפו ופסימ כו ערייים וופיי ביין ביין ביין ביין ביין ביין ביין ב	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete		Human dystrophin gene	601854981F1 NIH MGC 57 Homo sapiens culvia cidne liwa GE: 407 4546 5	602154364F1 NIH MGC_83 Home sapiens cUNA clone IMAGE: 4230034 3	IL5-HT0704-290600-108-c04 H10704 Homo sapiens culvA	Omithorhynchus anatinus coagulation factor X mKNA, comprete cos	Thermotoga maritima section 109 of 136 of the complete genuine	HSAAADTHS TEST1, Human adult 1estis tissue Homo sapiens clurk cione cam test2+*(5)	Human coagulation tactor VII (1-7) gene exch i and ractor VII (1-7	ne87f04.s.1 NCI_CGAP_Kid1 Home sapiens curve cione invaces i 1.200	yhesdo4.s1 Spares placenta Nozhir nomo saprars culta dulla d	QV4-NNC038-2/U400-18/-nuo NNV0050 nuito sapienio cutiva	Rattus norvegicus UDP-Gal:glucosylceramide beta-1,4-galactosyltransferase mRNA, complete cds	Homo sapiens mitochondrial glutathione reductase and cytosolic glutathione reductase (GRD1) gene,	complete cds, alternatively spliced	601338428F1 NIH_MGC_53 Homo saplens cDNA clone IMAGE:3680595 5	601338428F1 NIH_MGC_53 Homo sapiens cUNA cione IMAGE:3680995 5	yu07e10,r1 Soares fetal liver spleen 1NFLS Homo sapiens cLinA cione IMAGE 233130 3	Sus scrofa deoxyribonuciease II mRNA, complete cds	601452661F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3836398 3	Neisseria meningitidis DNA for region 2 (fhaB- and fhaC-homologs, unknown genes) and Hanking genes, strain FAM18	601140729F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3049830 5
-	Top Hit Database Source	EST HUMAN ele	MAN	HOMAN	1200	Т	EST HUMAN 90	NT cds				П	7	HUMAN			THUMAN		7	7	EST HUMAN Q	NT R	Ĭ				T_HUMAN		EST_HUMAN 6	N LN	EST_HUMAN 6
	Top Hit Acession No.	2 N99615.1				3.0E-02 BE889948.1		3.0E-02 AF213884.1		-		3.0E-02 BF246361.1					3.0E-02 Z21211.1			3.0E-02 R32019.1	AW895565.1	3.0E-02 AF048687.1		2.9E-02 AF228703.1	12 BE565644.1	2.9E-02 BE565644.1	H72805.1	12 AF060221.1	02 BF032233.1	AJ391284.1	2.9E-02 BE271437.1
	Most Similar (Top) Hit BLAST E Value	3.0E-02		3.0E-02	3.0E-02	3.0E-02	3.0E-02	3.0E-02		3.0E-02	3.0E-02	3.0E-02															2.9E-C	2.9E-C	2.9E-(
	Expression Signal	1.4		4.1	3.32	2.84	2.84	2.15		2.15	1.4	0.59	0.48	99.0	1.8	1.49				2	2.46	2.06		1.27				76.0			
	ORF SEQ ID NO:	31905		31906		32488	32489	32472			32885			34373		36221		37148						27891	L				31953		
	SEQ ID	18937		- [ı	ı	19467	19453		19453	19819	Ι.		L.		L.	_	L	L.	25389	24909	25383	1	15594	l	1	1	1	L	L_	1
	Probe SEQ ID NO:	6	3	6160	6692	9089	9089	6071		6971	7132	7483	8025	8539	8692	10357	10446	11197	11690	12243	12587	12629		2436	2890	2990	3908	5972	6199	6855	7148

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Table 4

		T	т-	_	7	_	7	_	_	_	_	_	-	_	_,	-	-	_		_	_:			_	<u>,</u>				_			
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Buchnera aphidicola natural-host Schlechtendalia chinensis gluconate-6-phosphate dehydrogenase (gnd) gene, partial cds	Buchnera aphidicola natural-host Schlechtendalla chinensis gluconata-6-phosphate dehydrogenase (gnd) gene, partial cds	CM3-PT0014-071299-051-c04 PT0014 Homo sapiens cDNA	CM3-PT0014-071299-051-c04 PT0014 Homo saniens cDNA	EST388706 MAGE resequences, MAGN Homo sapiens cDNA	Aeropyrum pernix genomic DNA, section 7/7	Sheep gene for ultra high-sulphur keratin protein	EST382234 MAGE reseguences, MAGK Homo seniens CDNA	Homo saplens retinal fascin (FSCN2) gene, expn 2	Homo saplens retinal fascin (FSCN2) gene exon 2	601594078F1 NIH MGC 9 Homo sapiens chiNA clone IMAGE 3048087 F	yd21b08.r1 Soares fetal liver splean 1NELS Homo seniens cDNA close MAGE: 40e982 21	Craterosticing plantacing m RNA for homeodomain lauring since protein (Ar. 1)	2898008.11 NCI CGAP GCB1 Home seniens cDNA close IMACE:771.1488 E	Cavia borcellus inwardiv-rectifikhor bobasetum channel Kirol Jiman competenti in anno 1900 de de de de de de de de de de de de de	Archaeoglobus fulgidus section 15 of 170 of the complete constants	V12h02.r1 Scares fetal liver spleen 1NFI S Homo sepiens CDNA Alone INAA CE 1. Joseph E1	Yeast CN31C chromosome III RAHS DNA (right arm transcription hot-spot)	Human germline T-cell receptor beta chain Dopamine-beta-hydroxylasse-like TRY1 TRY2 TRY3	TCRBV27S1P, TCRBV22S1A2N1T, TCRBV9S1A1T, TCRBV7S1A1N2T, TCRBV5S1A1T, TCRRV13S3	TCRBV6S7P, TCRBV7S3AZT, TCRBV13S2A1T, TCRBV9S2A2PT, TCRBV7S2A1N4T,	TCRBV13S9/13S>	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 6	yy86h12.r1 Soares_multiple_sclerosis_ZNbHMSP Homo septens cDNA clone iMAGE:786487 だ	y98h12.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE-280487 応	y/33d09.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:128657 6' similar to	SP:JC2284 JC2284 TISSUE FACTOR PATHWAY INHIBITOR - RHESUS	T.aestivum pTTH20 mRNA for wheat type V thionin	Oryza sativa mRNA for ascorbate oxidase, partial cds	A.bisporus pgika gene	ot96h03.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1624861 3'
gle Exon Prol	Top Hit Database Source	ĻΝ	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	Z.	TN	EST HUMAN	N	NT	EST HUMAN	EST HUMAN	NT	Т	N P	Т	HUMAN	Γ			•			Г	EST HUMAN		EST_HUMAN		NT	NT.	EST HUMAN
ig	Top Hit Acession No.	02 AF128279.1	02 AF129279.1	02 AW875979.1	02 AW875979.1	02 AW976597.1	02 AP000064.1	(55294.1	2.8E-02 AW970153.1		Г	Г	Γ	-	2.8E-02 AA280762.1						_			.2								
	Most Similar (Top) Hit BLAST E Value	2.9E-02	2.9E-02		2.9E-02	2.9E-02	2.9E-02	2.9E-02	2.8E-02 /	2.8E-02	2.8E-02/	2.8E-02	2.8E-02	2.8E-02	2.8E-02	2.8E-02	2.8E-02	2.8E-02 F	2.8E-02 X06322.1				· 2./E-02 U66059.1	2.7E-02 A	2.7E-02 N47268.1	2.7E-02 N		2.7E-02 R12245.1	2.7E-02 X61670.1	2.7E-02 AB004799.1	2.7E-02 X97580.1	2.7E-02 A
	Expression Signal	0.87	. 0.87	2.49	2.49	0.75	1.25	1.91	96'0	1.3	1.3	11.62	1.15	1.61	0.85	-	0.69	1.5	1.48			- {	3	1.74	1.92	1.92		1:2	0.86	0.64	0.93	2.29
	ORF SEQ ID NO:	33723		35396			36103	30590		28775	28776	30805	32670	34058	34749	34960	35058				~	20000	COSO7	28832	29645	29546		30841	31529	31612	1	32467
	Exon SEQ ID NO:	20592		22211			2289	17902	13335	16119	16119	18200	19626	20820	21606	21796	21891	25229	24876			2507	27	10.182	16918	16918	,	8012	200	18671	19270	19449
	Probe SEQ ID NO:	7897	7897	9558	8228	9774	10243	10977	662	3360	3360	5400	6711	8226	8915	9108	9212	12528	12630			4472	7/2	045 07 07 07 07 07	4178	4178	- 1	8	2812	883 2883	6505	6967

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Top Hit Descriptor	to:28g08.x1 Soares_total_fetus_Nb2HF8_9w Homo sepiens cDNA clone IMAGE:2065982.3' similar to contains Alu repetitive element;	transmembrane secretory component [human, leukocytes, Genomic, 657 nt, segment 4 of 11]	Homo sapiens chromosome 21 segment HS21 C082	ab02b02.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:839595 3'	Mus musculus histidine rich calcium binding protein (Hro), mRNA	Mus musculus histidine rich calcium binding protein (Hrc), mRNA	Mus musculus MHC class III region RD gene, partial cds; Bf, C2, G9A, NG22, G9, HSP70, HSP70, HSC70t, and smRNP genes, complete cds; G7A gene, partial cds; and unknown genes	Chicken dorsalin-1 mRNA, complete cds	Delnococcus radiodurans R1 section 151 of 229 of the complete chromosome 1	X852b04.X1 NCI_CGAP_Sar4 Homo sapiens cDNA clone IMAGE:2570383 3' similar to SW:Y069_HUMAN Q15041 HYPOTHETICAL PROTEIN KIAA0069 :	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63	qg27f11.x1 NCI_CGAP_Kid3 Homo saplens cDNA clone IMAGE:1762317 3'	801493473T1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3895578 3'	Vaccinla virus ORF1L, strain Wyeth	Vaccinia virus ORF1L, strain Wyeth	Rattus norvegious Nerve growth factor receptor, fast (Ngfr), mRNA	ak22f04.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1406719 3'	Homo sapiens KIAA1070 protein (KIAA1070), mRNA	Saccharomyces dairenensis NRRL Y-12639(T) ATP synthase subunit 9 (ATP9) gene, mitochondrial gene encoding mitochondrial protein, complete cds	Saccharomyces dairenensis NRRL Y-12639(T) ATP synthase subunit 9 (ATP9) gene, mitochondrial gene	encoding mitochondrial protein, complete cds	Homo sapiens chromosome 21 segment HS21C103	Homo sapiens radixin (RDX) mRNA	zs84c02.r1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:704162 5'	UI-HF-BN0-akj-e-10-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3077466 5'	602015501F1 NCI_CGAP_Brn64 Homo saplens cDNA clone IMAGE:4150944 5'	on28f08.y5 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1557827 5'	on26f06.y5 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1557827.5'	601680305R2 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950665 3
Top Hit Database Source	EST_HUMAN	NT	NT	EST_HUMAN			IN		LN	EST HUMAN	Т		EST_HUMAN	EST_HUMAN	NT	IN		EST_HUMAN		LΝ			NT			EST_HUMAN	EST_HUMAN	П	T	EST_HUMAN
Top Hit Acession No.	AI377036.1	S43442.1	AL163282.2	AA490021.1	6754241 NT	6764241 NT	A E 40000 R 4	L12032.1	AE002014.1	AW24154.1	AL161563.2	AL161563.2	AI206030.1	BE621748.1	Z99064.1	Z99064.1	TN 1721869	AA860946.1	11432020 NT	AF114952.1		AF114952.1	AL163303.2	4506468 NT	AA279351.1	AW500547.1	BF343827.1	AI793130.1	AI793130.1	BE974314.1
Most Similar (Top) Hit BLAST E Value	2.7E-02	2.7E-02	2.6E-02	2.6E-02	2.6E-02	2.6E-02	2 85 03		2.6E-02	2 RF.02		2.6E-02	2.6E-02	2.6E-02		2.6E-02	2.6E-02	2.8E-02	2.6E-02	2.6E-02	_			2.6E-02	2.6E-02	2.6E-02	2.6E-02	2.5E-02	2.5E-02	2.5E-02
Expression Signal	1.06	0.49	1.12	3.29	4.49	4.49	40	2.25	1.56	2.35	0.7	0.50	7.34	1.9	0.0	6.0	7.11	0.71	1.15	0.75		0.75	4.39	1.67	2.33	2.2	2.09	1.75	1.75	15.9
ORF SEQ ID NO:		34349			L				30332					32089		32594	32494	34232		35453	L		36165			37465	30615	25835		28224
Exon SEQ ID NO:	20946	1	ĺ	15088	15090	15090	4 E003		1	ſ		1_	Ĺ	19101	19562	19562	19471	21098		22267	_		22950	23856	24049	24152	25375	13303		13563
Probe SEQ ID NO:	8252	8514	558	2366	2368	2368	35.6	\$ 25	5005	5032	5754	2800	6125	6331	67.28	8728	6810	8403	9260	98 4		9614	10303	11191	11361	11563	12170	519	519	791

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		\neg	Т	Т	7	Т	÷	_	Т	т-	_	_	_	_	-	-	_		_	_	-	_			_	_	- ,	_	_	<u> </u>		_	
Single Exon Probes Expressed in Brain	Top Hit Descriptor	601980306R2 NIH MGC 83 Homo sapiens cDNA clone IMAGE 3950885 31	Rettus novegicus rabbhilin-3A mRNA, complete cds	H. carterae mRNA for fucoxanthin chlorophyll a/c binding protein Fcp1	H.carterae mRNA for fucoxanthin chlorophyl a/c binding protein Foot	PM2-NN0128-080700-001-a12 NN0128 Homo sapiens cDNA	PM2-NN0128-080700-001-a12 NN0128 Homo saplens cDNA	hf36h08.x1 Soares NFL T GBC S1 Homo sablens CDNA clone IMAGE: 3234018.3	zx83c10.x5 Soares overy tumor NbHOT Homo sepiens cDNA clone IMAGE:810354 3	7e30e09.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284008 3' similar to contains L1.t1 L1	601579393F1 NIH MGC 9 Homo sanlens CDNA clare IMAGE: 3029E4 F1	Chlamoromas rainhamhii VSB-3 m.BM. common sees sainhamhii VSB-3 m.BM. common sees sainhamhii VSB-3 m.BM.	602070662F1 NCI CGAP Brade Homo septems - DNA clare IMACE: 7212406 ET	602070562F1 NCI CGAP Rm84 Homo sanians CDNA clare IMACE 4233406 E	CHORDIN PRECURSOR (ORGANIZER-SPECIFIC SECRETED DORSAL JAING BACTOD)	wu08c10.x1 NCI CGAP GC6 Homo sapiens cDNA clone IMAGE-3518370 3	D.radicum 28S ribosomal RNA, D2 domain	qb22a08.x1 Soares pregnant uterus NbHPU Homo septens cDNA cione IMAGE·160e0p2 2	HYPOTHETICAL 46.7 KD PROTEIN C19G10.05 IN CHROMOSOMF I	HYPOTHETICAL 46.7 KD PROTEIN C19G10.05 IN CHROMOSOME I	Bos taurus partial stat6B gene, exons 17-19	Mus musculus major histocompatibility locus class II region: major histocompatibility protein class II alpha	chain (IAaipha) and major histocompatibility protein class II beta chain (IEbeta) genes, complete cds;	Homo septens dene for LECT2 complete cds	Homo seplens similar to AI FX3 protein (H. seplens) (1 OCRARA) DNA	Homo sablens mitorem-activated protein kinasa kinasa kinasa 13 /MA bak43)	601652365R2 NIH MGC 82 Homo saplens cDNA clone IMAGE:3835513 3	y75f11.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGF:211149 F	H-2 CLASS HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECLIBSOR (H-2K/R)	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K/B))	T.thermophila calcium-binding 25 kDa (TCBP 25) protein mRNA, complete cds	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))
gie Exon Pro	Top Hit Database Source	EST HUMAN	Į.	N.	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HIMAN	EST HUMAN	LN	EST HUMAN	EST HUMAN	SWISSPROT	EST HUMAN	N	EST HUMAN	SWISSPROT	SWISSPROT	TN		<u> </u>	L	Į	12	EST HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	NT		SWISSPROT
	Top Hit Acession No.	2.5E-02 BE974314.1	012571.1	X99697.1	X99697.1	02 BE701165.1	02 BE701165.1	02 AW 592114.1	02 AI732776.1	02 BE670128 1	02 BE746888.1	02 29029.1	02 BF526722.1	02 BF526722.1	02 091713	02 AW025821.1	02 X71303.1	02 Al147615.1	12 Q10335	12 Q10335	6.1			2.5E-02 AB007546.1	11420078		BE973327.1						
	Most Similar (Top) Hit BLAST E Value	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02		2.5F_02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.4E-02	2.4E-02	2.4E-02 P01901	2.4E-02	2.4E-02 P01901	2.4E-02
	Expression Signal	7.2	2.24	4.35	4.35	1	1	4.23	0.61	5.01	4.1	1,04	1.7	1.7	0.81	0.47	0.55	0.65	2.01	2.01	2:32		3.46	2.55	2.89	1.76	1.94	1.7	1.92	1.92	1.65	1.58	1.58
	ORF SEQ ID NO:	26289						29550	31335			31992	33336	33337	34680	34699		36356	36640	36641	36700		_		-	-	31043	27028	27501	27502	29702	29847	29848
	Exon SEQ ID NO:	Н	15471						18422	18878	18893	19018	20233	20233	21418	21554	22614	23128	23401	23401	23456		23478	24361	25311	25182	24804	14339	15584	16584	17074	17220	17220
	Probe SEQ ID NO:	840	2766	2956	2956	4023	4023	4182	5625	6100	6115	6244	7563	7563	8724	8863	9966	10482	10712	10712	10773		10795	11770	12134	12311	12432	1593	2037	2037	4335	4485	4485

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit / N	Top Hit Database Source	Top Hit Descriptor
2068	17787	30403	0.95	2.4E-02	8922702 NT	INT	Homo sapiens hypothetical protein FLJ10844 (FLJ10844), mRNA
6121	18899		6.0	2.4E-02	2.4E-02 W86680.1	EST_HUMAN	zh63h04.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:416791 3'
6267	19040			2.4E-02	2.4E-02 M31650.1	NT	Chicken myristoylated alanine-rich Cikinase substrate (MARCKS) mRNA, complete cds
6267	19040		0.58	2.4E-02	2.4E-02 M31650.1	۲N	Chicken myrtstoylated alanine-rich C kinase substrate (MARCKS) mRNA, complete cds
7121	19809	32875	0.8	24E-02	Z20573.1	EST_HUMAN	HSAAACKVX T, Human adult Rhabdomyosarcoma celHine Homo sapiens cDNA
7138	ļ			2.4E-02	2.4E-02 X12925.1		Rat gene for uncoupling protein (UCP)
7138	19825		6.0	2.4E-02	2.4E-02 X12925.1	TN	Rat gene for uncoupling protein (UCP)
7791	20486			2.4E-02	2.4E-02 AW813007.1	EST HUMAN	RC3-ST0186-230300-019-h06 ST0186 Homo sapiens cDNA
7844	1		0.5	2.4E-02	2.4E-02 M16780.1	TN	Human retrotransposon 3' long terminal repeat
8340	21033		09 0	2 4F.02	2 HZR3ZR 1	FST HIMAN	yu12c05.s1 Soares fetal liver spleen 1NFLS Homo sepiens cDNA clone IMAGE:233576 3' similar to contains. Alu repetitive element contains A3R repetitive element:
							2835611.s1 Soares fetal liver spleen 1NFLS Homo sepiens cDNA clone IMAGE:2945963' similar to
8429						EST_HUMAN	gb K02909 RATSR7K Rat (rRNA);contains A3R.b1 A3R repetitive element;
8885	21576	34718	0.57		2.4E-02 AE001125.1	TN	Borrella burgdorferi (section 11 of 70) of the complete genome
	ı						zu91c06.s1 Soares_lestis_NHT Homo sapiens dDNA clone IMAGE:745354 3' similar to gb:J04422 ISLET AMY OID POI YPEPTIDE PRECI IRSOR (HI IMAN) contains Alu renetitive element contains element XTR
8907	21598	34740	0.78	2.4E-02	2.4E-02 AA625660.1	EST_HUMAN	XTR repetitive element;
9591	1				2.4E-02 AF124160.1	N	Arabidopsis thaliana molybdopterin synthase sulphurylase (cnx5) gene, complete cds
9591	22244	35428			AF124160.1	뉟	Arabidopsis thaliana motybdopterin synthese sulphurylase (cnx5) gene, complete cds
9206	1		2.38		2.4E-02 AV692954.1	EST_HUMAN	AV692954 GKC Homo saplens cDNA clone GKCDSC03 6'
	1	<u>.</u>					nh07b12.s1 NCI_CGAP_Thy1 Homo sepiens cDNA clone IMAGE:943583 similar to contains Alu repetitive
988	22331	35/28	2.73	١	2.4E-02 AA493894.1	ES HOMAIN	General contents defined in 1 No legetive defined;
10512	23158		0.46	1	BE38/111.1	ESI HUMAN	OF 124 BOZET I NITE MODE AND SEPTEMBERS COME INVACES SO I BOZET IN THE MODE OF THE SOURCE SO I BOZET IN THE SOURCE
11565	24164	37475		2.4E-02	2.4E-02 AF109805.1	Ę	Mus musculus major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, CLCP, NG24, NG25, and NG28 genes, complete cds; and unknown genes
		1					
11565		37476	1.89	2.4E-02	2.4E-02 AF109905.1	Ł	Mus musculus major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, CLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes
11938	24495			2.4E-02	TN 60627906	FX	Bacteriophage blL67, complete genome
12081	24589			2.4E-02	6753635 NT	INT	Mus musculus DinB homolog 1 (E. coli) (Dinb1), mRNA
12136	24625	31094	2.37	2.4E-02	2.4E-02 BE928869.1	EST_HUMAN	MR0-FT0175-310800-202-a06 FT0175 Homo sapiens cDNA
12186	24657	31063	1.66	2.4E-02	2.4E-02)U78167.1	NT	Rattus norvegicus cAMP-regulated guanine nucleotide exchange fector I (cAMP-GEFI) mRNA, complete cds

Page 156 of 536 Table 4

| Top Hit Descriptor | Raffus nervectors cAMP remitted months | Home sapiens SNCA isoform (SNCA) pana complete od the sapiens of the sapiens SNCA isoform (SNCA) was complete od the sapiens SNCA isoform (SNCA) was complete od the sapiens SNCA isoform (SNCA) was complete of the sapiens SNCA isoform (SNCA) was complete of the sapiens SNCA isoform (SNCA) was complete of the sapiens of t | Caenorhabditis elegans mRNA for Iron-sulfur subunit of mitochondrial succinate dehydrogenase, complete | 2384408 of Same detail to the little details | 4 Homo seniens memmen it most consolidation in 195 August Clone IMAGE:299294 5: | Scenarista chomosome IV results from OBEVEL AND AND | HSAAACADH P Himen footal Brain What it | Can's befactalarhaides hadran potential (OA) (O3) | Gallus gallus consolar 456 (PARS) and | General School Connext 45.6 (CARA R) con complete cas | CM4-NNORD-200401-150-NORD Manager
Manager Mana | CM3-MT0118-010900-318-007 MT0118 Home series - DNIA | CM34AT0418-040900-348-07 ArT0418 Home con Child | XS25408 X1 NCL CCAP TIP Home conformation and Alberta County

 | XS2BG08 Y NCI COAP 112 Home capture COM LINE COAP 13 | 601672279F1 NIH MGC 20 Home septemble CONA size 1440 F120 F1

 | 601672279F1 NIH MGC 20 Homo sapiens CDNA claire IMAGE:3958386 5 | Caulobacter crescentus topolsomerase IV ParE subunit (parE) gene, complete cds, and propionyl-CoA

 | Arabidoosis thaliana DNA chromosomo 4 contra cas | MR0-HT0080-011099-002-009 HTD080-HTM0-17 | Human plactin (PLFC1) game aware 3.32 and complete 1.1 | Homo sablans PDX1 gene for thoughout comments V | Homo saplens DDX1 nane for linear contribution of the contribution | wa78h10 v1 Scenes NEI T CBC of Little
 | wa76h10 x1 Soares NFT T GRC 34 Home explens cDNA clone IMAGE;2302147 3 | HYPOTHETICAL 35.6 KD PROTEIN BROSE & IN CUBOLICE AND THE BROSE AT 11 BRITAINS | CHROMOSOME ASSEMBLY PROTEIN XOAP.C | Scherichia coli K-12 MG1655 section 89 of 400 of the complete concess | Escherichia coli K-12 MG1635 section 89 of 400 of the complete genome | Bacillus licheniformis isolate N57N1 KerA gene, partial cds
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| Top Hit Acession
No. | U78167.1 | AF163864.1 | AB008569 1 | W05340.1 | U94165.1 | 274283.1 | 220377.1 | 23429.1 | | | AW899107.1
 | 3E93525.1 | 3E835225.1 |

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Value | 2.4E-02 | 2.4E-02 | 2.4E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02 | 2.3E-02
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 | 2.3E-02 E | 2.3E-02.L

 | 2.3E-02 | 2.3E-02 B | 2.3E-02 ∪ | 2.3E-02 A | 2.3E-02 A | 23E-02 A
 | 2.3E-02 A | 23E-02 P | 2.3E-02 P | 2.3E-02 A | 2.3E-02 A | 2.3E-02 AF282894.1
 |
| Expression
Signal | 1.66 | 1.34 | 3.88 | 4.20 | 10.45 | 2.08 | 6.19 | 8.0 | 1.06 | 1.06 | 0.93
 | 0.88 | 0.88 | 1.14

 | 1.14 | 2.58

 | 2.58 | 3.63

 | 4.08 | 0.69 | 6.28 | 0.94 | 0.94 | 0.68
 | 0.68 | 0.98 | 0.77 | 1.33 | 1.33 | 0.46
 |
| ORF SEQ
ID NO: | 31104 | | | | | 27809 | 29084 | | 29499 | 29500 | 29755
 | 29780 | 29781 | 29782

 | 29783 | 29919

 | 29920 | 30756

 | 32292 | 30513 | 33595 | 34204 | 34205 | 34429
 | 34430 | 34880 | 35617 | 35776 | 35777 | 36397
 |
| | | _ | 24764 | 14603 | 14817 | - 1 | - 1 | | | | - 1
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 | - 1 | 17290

 | 17290 | 18096

 | 19288 | 17959 | 20472 | 21083 | 21063 | 21289
 | 21289 | 21726 | 22410 | 22577 | 22577 | 23170
 |
| Probe
SEQ ID
NO: | 12186 | 12218 | 12360 | 1865 | 1880 | 2350 | 3670 | 3702 | 4129 | 4129 | 4386
 | 4416 | 4415 | 4416

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19 | 4555

 | 4555 | 5291

 | 6522 | 9883 | 7778 | 8370 | 8370 | 8597
 | 8597 | 8038 | 9769 | 8266 | 8838 | 10524
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Source 24657 31104 1.86 2.4E-02 U78167.1 NT 24784 3.88 2.4E-02 AF163864.1 NT 14603 4.29 2.3E-02 W05340.1 EST HUMAN 14603 4.29 2.3E-02 W05340.1 EST HUMAN 16412 2.28094 6.19 2.3E-02 U94165.1 NT 16871 2.9094 6.19 2.3E-02 L24293.1 NT 16871 2.9499 1.06 2.3E-02 L24293.1 NT 16871 2.9600 1.06 2.3E-02 L24299.1 NT 171723 29769 1.06 2.3E-02 L24799.1 NT 17162 29776 0.63 2.3E-02 L24799.1 NT 17162 29776 0.08 2.3E-02 R24896907.1 EST HUMAN 17162 29778 0.08 2.3E-02 R2608693693.1 EST HUMAN 17890 29763 <td< td=""><td>Exam
NO: ORF SEQ
ID NO: Expression
Signal (Top) Hit
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Source 24678 31104 1.86 2.4E-02 U78167.1 NT 24784 3.88 2.4E-02 AF163864.1 NT 14603 4.29 2.3E-02 W05340.1 EST HUMAN 14607 2.7809 2.06 2.3E-02 W05340.1 EST HUMAN 16472 2.7809 2.06 2.3E-02 W05340.1 EST HUMAN 16473 2.3E-02 W05340.1 EST HUMAN NT 16450 0.88 2.3E-02 L24263.1 NT 16871 2.3604 0.08 2.3E-02 L24269.1 NT 16871 2.3609 1.06 2.3E-02 L24789.1 NT 171723 29765 0.83 2.3E-02 L24789.1 NT 17162 29781 0.88 2.3E-02 R2489.07.1 EST HUMAN 17162 29782 0.88 2.3E-02 R2489.1 EST HUMAN 1780 29782 1.14 <</td><td>Exon
NO: ORF SEQ
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Signal (Top) Hit
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NO: ORF SEQ
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NO: ORF SEQ
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NO: ORF SEQ
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NO: ORF SEQ
Signal Expression
Signal Most Similar
(Top) Hit
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Value Top Hit Acession
Source 24678 31104 1.86 2.4E-02 AE-02 AF-163864.1 NT NT 14673 2780 2.3E-02 AF-163864.1 NT NT NT 14673 2.780 2.3E-02 AF-163864.1 NT NT NT 14673 2.780 2.3E-02 AF-16386.1 NT NT NT 16473 2.780 2.3E-02 AF-1638.1 NT NT NT 16473 2.780 0.8 2.3E-02 AF-1748.1 NT NT 16473 2.780 0.8 2.3E-02 AF-1748.1 NT NT 17162 2.780 0.8 2.3E-02 AF-1748.1 NT NT 17280<</td><td>Exam ORF SEQ Expression (Top) Hit Acession Top Hit Acession
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NO: ORF SEQ
ID NO: Expression
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Source 24678 31104 1.86 2.4E-02 U78167.1 NT 24784 3.88 2.4E-02 AF163864.1 NT 14603 4.29 2.3E-02 W05340.1 EST HUMAN 14607 2.7809 2.06 2.3E-02 W05340.1 EST HUMAN 16472 2.7809 2.06 2.3E-02 W05340.1 EST HUMAN 16473 2.3E-02 W05340.1 EST HUMAN NT 16450 0.88 2.3E-02 L24263.1 NT 16871 2.3604 0.08 2.3E-02 L24269.1 NT 16871 2.3609 1.06 2.3E-02 L24789.1 NT 171723 29765 0.83 2.3E-02 L24789.1 NT 17162 29781 0.88 2.3E-02 R2489.07.1 EST HUMAN 17162 29782 0.88 2.3E-02 R2489.1 EST HUMAN 1780 29782 1.14 < | Exon
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NO: ORF SEQ
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Source 24657 31104 1.86 2.4E-02 U78167.1 NT 24678 31104 1.86 2.4E-02 U78167.1 NT 24764 3.88 2.4E-02 U78167.1 NT 14603 2.7809 2.0E 2.3E-02 U94168.1 NT 16423 2.9064 6.19 2.3E-02 U94168.1 NT 16425 2.9499 1.06 2.3E-02 L24728.1 NT 164871 2.9499 1.06 2.3E-02 L24728.1 NT 164871 2.9499 1.06 2.3E-02 L24728.1 NT 17162 2.9780 0.93 2.3E-02 L8695225.1 EST HUMAN 17780 2.9781 0.88 2.3E-02 BE935225.1 EST HUMAN 17290 2.9780 0.68 2.3E-02 BE925225.1 EST HUMAN <</td><td>Exon
NO: ORF SEQ
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2.4E-02 ID Hit
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Patabase 24657 31104 1.86 2.4E-02 ID 18467.1 NT 24678 3.88 2.4E-02 ID 18467.1 NT 14603 4.29 2.3E-02 ID 18465.1 NT 14677 2.064 6.19 2.3E-02 ID 18465.1 NT 1672 2.7809 2.08 2.3E-02 ID 2423.1 NT 16453 2.9604 6.19 2.3E-02 ID 2423.1 NT 16472 2.7809 0.8 2.3E-02 ID 2423.1 NT 16455 2.9604 6.19 2.3E-02 ID 2423.1 NT 1647 2.9499 1.06 2.3E-02 ID 2423.1 NT 1712 29780 0.89 2.3E-02 ID 2429.1 NT 1716 2.9781 0.88 2.3E-02 ID 2429.1 NT 17780 29780 0.08 2.3E-02 ID 24083.1 EST HUMAN 17780 29878 1.14 2.3E-02 ID 24083.</td><td>Exon
NO: ORF SEQ
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Source 24657 3104 1.86 2.4E-02 U78467.1 NT 24678 3104 1.86 2.4E-02 U78467.1 NT 24678 3.88 2.4E-02 U78467.1 NT 14603 4.29 2.3E-02 U8465.1 NT 16072 2.7809 2.08 2.3E-02 U8465.1 NT 16473 2.9604 6.19 2.3E-02 U8465.1 NT 16474 2.9604 6.19 2.3E-02 U8405.1 NT 16475 2.9780 0.8 2.3E-02 U8405.1 NT 16871 2.9800 1.06 2.3E-02 L24799.1 NT 17122 29780 0.83 2.3E-02 L24799.1 NT 17162 29781 0.88 2.3E-02 L24799.1 NT 17780 29782 1.14 2.3E-02 L24799.1 NT
 17780 29781 1.14 2.3E-02 L24799.1 NT 1</td><td>Exon
NO: ORF SEQ
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Top) Hit
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(Top) Hit
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No: Top Hit Acession
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Source Top Hit Acession
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Top) Hit
ID NO: Top Hit Acession
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Patabase 24657 31104 1.86 2.4E-02 ID 18467.1 NT 24678 3.88 2.4E-02 ID 18467.1 NT 14603 4.29 2.3E-02 ID 18465.1 NT 14677 2.064 6.19 2.3E-02 ID 18465.1 NT 1672 2.7809 2.08 2.3E-02 ID 2423.1 NT 16453 2.9604 6.19 2.3E-02 ID 2423.1 NT 16472 2.7809 0.8 2.3E-02 ID 2423.1 NT 16455 2.9604 6.19 2.3E-02 ID 2423.1 NT 1647 2.9499 1.06 2.3E-02 ID 2423.1 NT 1712 29780 0.89 2.3E-02 ID 2429.1 NT 1716 2.9781 0.88 2.3E-02 ID 2429.1 NT 17780 29780 0.08 2.3E-02 ID 24083.1 EST HUMAN 17780 29878 1.14 2.3E-02 ID 24083. | Exon
NO: ORF SEQ
Signal Expression
Signal (Top) Hit
I (Top) Hit
Top Hit Acession Top Hit
Acession
No. Top Hit
Source 24657 3104 1.86 2.4E-02 U78467.1 NT 24678 3104 1.86 2.4E-02 U78467.1 NT 24678 3.88 2.4E-02 U78467.1 NT 14603 4.29 2.3E-02 U8465.1 NT 16072 2.7809 2.08 2.3E-02 U8465.1 NT 16473 2.9604 6.19 2.3E-02 U8465.1 NT 16474 2.9604 6.19 2.3E-02 U8405.1 NT 16475 2.9780 0.8 2.3E-02 U8405.1 NT 16871 2.9800 1.06 2.3E-02 L24799.1 NT 17122 29780 0.83 2.3E-02 L24799.1 NT 17162 29781 0.88 2.3E-02 L24799.1 NT 17780 29782 1.14 2.3E-02 L24799.1 NT 17780 29781 1.14 2.3E-02 L24799.1 NT 1 | Exon
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(Top) Hit
PLASTE Top Hit Acession
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Source 24678 31104 1.86 2.4E-02 AE-02 AF-163864.1 NT NT 14673 2780 2.3E-02 AF-163864.1 NT NT NT 14673 2.780 2.3E-02 AF-163864.1 NT NT NT 14673 2.780 2.3E-02 AF-16386.1 NT NT NT 16473 2.780 2.3E-02 AF-1638.1 NT NT NT 16473 2.780 0.8 2.3E-02 AF-1748.1 NT NT 16473 2.780 0.8 2.3E-02 AF-1748.1 NT NT 17162 2.780 0.8 2.3E-02 AF-1748.1 NT NT 17280< | Exam ORF SEQ Expression (Top) Hit Acession Top Hit | Exameration Corporation
Page 157 of 536 Table 4 Single Exon Probes Expressed in Brain

ne47h07.s1 NCI_CGAP_Cc3 Homo sepiens cDNA clone IMAGE:900541 3' similar to contains Alu repetitive GLUCOAMMLASE S1/S2 PREGURSOR (GLUCAN 1,4-ALPHA-GLUCOSIDASE) (1,4-ALPHA-D-GLUCAN Dictyostelium discoideum extracellular signal-regulated protein kinase (ERK1) mRNA, complete cds Homo sapiens dead ringer (Drosophila)-like 1 (DRIL1), mRNA Columba livia nucleoside diphosphate kinase (NDPK) gene, nuclear gene encoding mitochondrial protein, Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, S.pneumoniae popA gene and open reading frames nn24a04.s1 NCI_CGAP_Gas1 Homo sapiens dDNA clone IMAGE:1084782.3 Homo sapiens chromodomain helicase DNA binding protein 2 (GHD2) mRNA MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK) Infectious bursal disease virus segment B strain IL4 VP1 gene, complete cds PM0-BT0340-170100-004-b03 BT0340 Homo sapiens cDNA Mus musculus Sjogren syndrome antigen A1 (Ssa1), mRNA 601584309F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3938571 5 AV761502 MDS Homo sapiens cDNA clone MDSADG01 5' Dictyostellum discoldeum histidine kinase C (dhkC) mRNA, complete cds 601179958F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3546567 Streptomyces sp. alpha-1,3/4 fucosidase precursor gene, complete cds Arabidopsis thallana DNA chromosome 4, contig fragment No. 27 MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK) Arabidopsis thaliana DNA chromosome 4, contig fragment No. Top Hit Descriptor S.cerevisiae chromosome XVI reading frame ORF YPL241c AV699721 GKB Homo sapiens cDNA clone GKBAND03 3' nembrane protein 1 (TMEM1), mRNA S.cerevisiae chromosome IV reading frame ORF YDL 2454 P. vulgata alpha tub 2 mRNA GLUCOHYDROLASE) complete cds) complete cds) complete cds element; EST_HUMAN EST_HUMAN NT EST_HUMAN EST HUMAN EST_HUMAN HUMAN EST_HUMAN SWISSPROT SWISSPROT Top Hit Database Source 7.1 NT 11426388 NT 눋 ż Ż 'n È 6678140 11423632 4557448 Top Hit Acession 2.2E-02 AB026898.1 2.2E-02 BE797601.1 2.2E-02 H 114230 2.2E-02 AA503553.1 2.2E-02 P07313 2.2E-02 P07313 2.2E-02 Z82001.1 2.2E-02 AA577785.1 2.2E-02 AF083094.1 2.1E-02 AV761502.1 2.1E-02 AF029726.1 AV761502.1 2.2E-02 AB026898.1 AV699721.1 2.2E-02 AL 161515.2 2.2E-02 AL161515.2 2.2E-02 AF018267.1 2.2E-02 45 2.3E-02 BE278331.1 ģ 2.2E-02 Z73597.1 2.2E-02 AV699721 X79468.1 Z74293.1 2.3E-02 U39394.1 2.3E-02 U11077.1 P08640 2.3E-02 (Top) Hit BLAST E **Jost Similar** 9.98 4.07 6.11 1.66 1.54 1.54 2.22 1.28 1.38 90. 0.75 2.16 5.07 2.42 162 4.13 Expression Signal 37115 29221 29290 32902 35422 35423 34104 34105 34546 36615 30997 26147 27469 ORF SEQ ΩÖ 13195 22239 22239 24737 13222 20963 20963 21401 23834 24425 23373 25260 14743 16585 13494 16390 14497 14497 SEQ ID 24892 ÿ 12315 436 3834 3834 3899 5006 7146 9586 11841 410 10105 11167 12616 174 9286 Probe SEQ ID 12058 1755 2008 3428 8269 8269 8709 12562 1755 ö

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Table 4
Single Exon Probes Fyunsson

Single Exon Probes Expressed in Brain	Most Similar (Top) Hit Acession (Top) Hit Dalabase Signal BLASTE No. Source	10.32 2.1E-02 U72073.1 NT Complete and South Cott (cott), Cott (cott.), and spore cost protein Cottin (cottin) genes.	32 AF204395.1 NT	2 AF204395.1 NT	02 P02438	2.1E-02 P02438 SWISSPROT	2.1E-02 P02438 SWISSPROT	2.1E-02 BE072546.1 EST HUMAN	2.1E-02 BE072546.1 EST HUMAN	2.1E-02 AA225095.1 EST HUMAN	2.1E-02 N29266.1 EST HUMAN	2 BE072546.1 EST HUMAN	T	EST HUMAN	2 274293.1	2.1E-02 BF343656.1 EST HUMAN	2.1E-02 U44914.1	2.1E-02 AI768127,1 EST HUMAN	2.1E-02 V19213.1		2.1E-02 AL163302.2 NT	EST HUMAN	2.1E-02 AW378529.1 EST HUMAN	2.1E-02 BF086199.1 EST_HUMAN	238 NT		2.1E-02 A 1243243 4 NT	2.1E-02 A.1243213.1		1.15) 2.1E-02 L28324.1 NT UmuD MucA homolog genes, complete cds; and unknown genes	2.1E-02	AP001510 4 NIT
	単 出 山		2.1E-02 AF	2.1E-02 AF	2.1E-02 PO	2.1E-02 P0;	2.1E-02 P0	2.1E-02 BE	2.1E-02 BE(2.1E-02 AA	2.1E-02 N28	2.1E-02 BEC	2.1E-02 BE0	2.1E-02 AA4	2.1E-02 Z74	2.1E-02 BF3	2.1E-02 U44	2.1E-02 AI76	2.1E-02 Y19	2.1E-02 Y08	2.1E-02 AL1	2.1E-02 AI82	2.1E-02 AW	2.1E-02 BF0	2.1E-02					=+	2.1E-02 AA98	2 1E OSTABOO
	Expression Signal	10.32	1.21	1.21	1.06	1.06	1.06	1.2	1.2	1.32	4.48	1.07	1.07	1	0.81	0.81	1.47	1.53	0.69	4.51	1.05	0.76	1.13	0.88	9.0	25	2.61	2.61		2	0.69	7 V
	ORF SEQ ID NO:	26656				27219	27220	27492	27493	28041	26193	27492	27493	28973	29480	29641	29775	29788		30031	30122	30124	31259	32466	34249	35236	35368	35369	00130	25/30	35814	36305
	Exon SEQ ID NO:		- 1	ŀ		- !	1		- 1	- 1	- 1	14763	14763	16326	16853	17014	17147	17157	17197	17396	17494	17501	18350	28 48 8	21110	22065	22184	22184	69.00	3	22609	22168
	Probe SEQ ID NO:	1240	1366	1366	1775	1775	1775	2028	2028	2591	2819	3147	3147	3671	<u>\$</u>	4275	<u>\$</u>	4421	4461	882 283	4762	4769	5553	888	\ <u>\</u>	9403	9531	9531	6000	3	9961	10522

Page 159 of 536 Table 4 Single Exon Probes Expressed in Brain

Too Lif Descriptor		Mus musculus heat shock protein, 74 kDa, A (Hspaßa), mRNA	Homo sapiens putative psinHbA pseudogene for flati Kerauli, exclusion and managed 1.22	Human germline UBE-1L gene similar to the gene for uprquium-acusating enzyme, control and	Azospirillum brasilense major outer membrane protein OmaA precursor (omaA) gene, complete cds	7g51c08.x1 NCI_CGAP_Pr28 Homo capiens cDNA clone IMAGE:3309998 3 similar to contains intervised. MER1 repetitive element ;	QV4-NN0038-270400-187-h05 NN0038 Homo sapiens cDNA	Mus musculus DinB homokog 1 (E. coli) (Dinb1), mRNA	aa15b10,r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:813307 5	Mus musculus DinB homolog 1 (E. coli) (Dinb1), mKNA	Homo sepiens genomic region containing hypervariable minisatellites chromosome 1 (1950,33) or nonio	Septembrane Septembrane Septembrane (F. 140379) mRNA	Homo saptens hypothetical protein F1 10070 (F1 10070) mRNA	Homo sapiens nypoureucal protein 1 to 1007 9 (to 100.00) mmm.	omo sapiens hypothetical protein FLJ 10469 (FLJ 10468), minno	Homo sepiens hypothetical protein run 10480 (Turi 10480), illinoin	Arabidopsis thallana DNA chromosome 4, contg fragment No. 32	7g51c08.x1 NC_CGAP_P728 Homo sapiens cDNA cidne Introductions of pilling to contain the contained that the contained the contained that the contained the contained that the contained t	Mus musculus sema domain, transmembrane domain (TM), and cytopiasmic domain, (semaphorin) 6B	(Seme6b), mRNA	Arabidopsis thallana C2H2 zinc finger protein F-ZF mKNA, complete cus	P. vulgaris hydroxyproline-rich glycoprotein (HRGP) mrvvA, 3 et la	Caenorhabditis elegans sma-z mrNA, complete cus	Dictyostelium discoideum class VII unconventional mycsm (myor) gene, curipred cus	Pyrococcus horikoshii OT3 genomic DNA, / / / / UOT-984/UU nt. posludii (#7)	Pyrococcus horikoshii OT3 genamic DNA, 777001-994000 nt. posiuon (**/)	Japanese encephalitis virus envelope protein mKNA, paruai cos	wa17502.x1 NCI_CGAP_Kid11 Homo sapans conversion to the livings.	Mycobacterium tuberculosis H3/RV compreis gendlie, sogilitari sociationalismosta.	Equus cabalus UNA (v) 1/alphanityu kwyasod (1, ko-1)kwo; on the control of mana (1, mRNA	Homo sapiens annymin 3, more of Ranvier (ankvrin G) (ANK3), transcript variant 1, mRNA	
Top Hit	Database Source		Ĭ	I L	-A	7 M NAMINAN M	T		T_HUMAN								N⊤	FST HUMAN	Т		NT				NT.			T HUMAN				
Too Hit Acession	Ŋ.	6754255 NT	Y19213.1		2 AF183913.1		2.0E-02 Br002832.1	3635		2.0E-02 6753635 NT		2.0E-02 AL096805.1	8922391 NT	8922391 NT	8922453 NT	8922453 NT	2.0E-02 AL161532.2	2 OF 02 BE003033 1	DI 002602. I	7305474 NT	02 AF095588.1	2.0E-02 M18095.1	U34778.1	02 135321.2	32 AP000004.1	02 AP000004.1	02 U70408.1	02 AI640342.1	02 Z73966.1	D88184		10947055IN
Most Similar		2.1E-02	2.1E-02	2.1É-02 L34170.1	2.1E-02/	200	20E-02	205.02	2.0E-02/	2.0E-02		2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	20 0	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-02	2.0E-	2.0E-	2.0E-02	2.0E-02	2.0E-02
1	Signal	1.38	8.62	1.89	5.71	;	1.1	2 76	2.72	2.11		1.6	1.17	1.17	2.39	2.39	3.24		2.11	4:	2.35	1.3	0.58	0.7	1.11					2.17		2.21
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Exon	SEQ ID	24080	17197	25163	24080		12844	1	13001	13553	3	13823	13930	13930	L	L			12844	15905			L	1_	L	1		1_	1	L	24277	24277
Probe	SEQ ID	11479	12206	12339	40774	2	9 ;		707	3 2	•	1065	1177	1177	1866	1866	2891		3077	2444	322	3988	5548	6807	245	7,450		10259	10539	11344	11682	11682

Page 160 of 536 Table 4

313307 F	0 %	is in the second	o contains L1.t1 L1								331 3'	ins Alu repetitive					1 3' similar to														
aa15b10,r1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGF	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 32	yd04c09.r1 Soares Infant brain 1NJB Homo sapiens cDNA clone IMAGE: 24875	hf19a07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:914196 similar to contains L1.t1 L1	EMPTY SPIRACI ES HOMEOTIC BROTEIN	Homo saniene chronosomo 24	Homo saniens chammens 24	Arabidoosis theirms DNA chemical and the control of	modefield Inc. Code Sea U.	AVEARES GIT Home carried appears clow cone IMAGE: 1238337 3'	228603 of Section September CONA GOOD GLOBILHO7 3	601572882F1 NIH MGC 57 Home scales CNA 111 A Clone IMAGE:284	qn04c07.x1 NO_CGAP_Lu5 Homo saplens cDNA clone IMAGE:1897280 3' similar to contains Alu remetitive	element;	Mycoplasma Imitans VIhA1 precursor (whA1) and VIhA2 precursor (whA2) genes, partial cds	HOMEO IIC BICOID PROTEIN (PRD-4)	HOMEOTIC BICOID PROTEIN (PRD-4)	yeouve.x1 soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE.214466	Arabidonsis thelians DNA shares	Vius musculus T cell receptor gamma (Am. TOB	Veleagns gallopavo paraoxonasa-2 (PON2) mBNA America danma 3 gene clusters	Prosophila kanakoi gene for alvami Anhambai A. A.	Tomo sapiens interferon damma recentor alpha chair anno 1	lomo sablens interferon commo coccess. I	deisseria maninalitie serveren. A designation of the serverence of	101896130F1 NIH MGC 10 Home serial AZ491 complete genome; segment 3/7	licoligna tabasalim fire 1 - h. f. f. f. f. f. f. f. f. f. f. f. f. f.	01852388F1 NIH MCC se U	Mechanistis sp PCC8803	Torio cholerae V86 nbs. 2 minute garome, 20/2/, 2639000-2644764	Irudo medicinalis intermediata filament dilarin mBNA complete cds	Candida albicans lambda Ca3/B fragment
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1.9	1.82	4.0	2.15	1.15	2.68	2.68	76.0	7.48	1.66	1.18	9.58	98	1 49	183	188		3.21	4.09	0.99	1.41	0.86	1.38	1.38	1.23	1.03	9.0	1.05	0.54	1.44	2.82	1.36
cacos]		26095	27047	27496	27497	27968	28320	28369			29095	29402	29539	29540		29872	27968	30667	30872		32744	32745		35082	35452	35798	36017	36614	30903	-
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14767 2749F 2.6B 1.9E-02 P18488 NT 1627 2749F 2.6B 1.9E-02 P18488 NT 1627 2749F 2.6B 1.9E-02 P18388 NT 1657 2789B 0.97 1.9E-02 P18383 NT 1657 2780B 0.97 1.9E-02 P184660.2 NT 1667 2780B 0.97 1.9E-02 P41163361 EST_HUMAN 1671B 28369 1.9E-02 P479486631 EST_HUMAN	15006 1.82 2.0E-02 AA456638.1 EST_HUMAN 15606 1.82 2.0E-02 AL161532.2 NT_EHUMAN 13452 25096 2.15 1.9E-02 P18488 SWISSPROT 14767 2.7496 2.68 1.9E-02 P18488 SWISSPROT 14767 2.7496 2.68 1.9E-02 AL163303.2 NT_EHUMAN 1527 2.89309 7.48 1.9E-02 AL163503.2 NT_EHUMAN 15718 2.8320 7.48 1.9E-02 AL163669.1 EST_HUMAN 15718 2.83599 1.66 1.9E-02 AR133669.1 EST_HUMAN 15718 2.83599 1.66 1.9E-02 AR33669.1 EST_HUMAN 1.8E-02 AR33699 AR3	17805 178 2.0E-02 AA466538.1 EST_HUMAN 26036 1.82 2.0E-02 AL161532.2 NT_HUMAN 25038 6.4 2.0E-02 T80037.1 EST_HUMAN 13452 25095 2.15 1.9E-02 P4848 SWISSPROT 14767 27496 2.68 1.9E-02 P418333.2 NT 14767 27497 2.68 1.9E-02 P4163303.2 NT 15572 27808 0.97 1.9E-02 P4163603.2 NT 15672 28320 7.48 1.9E-02 P4163660.1 EST_HUMAN 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1.95-02 A.1616				

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	Top Hit Descriptor	hn52c06.x1 NCI_CGAP_Co17 Homo saplens cDNA clone IMAGE:3027274 3' similar to contains element. MER29 repetitive element;	89452991 NIT MGC 17 TOTO SEPTION COLOR COL	H.francisci mKNA for myelin basic protein (wibr.)	Drosophila melanogaster cyuphashiic procein endo (u.v.) iiii s.v., ciiii.	Pseudomonas aeruginosa FAU1, securor 195 or u.s. or u.	1922/2019/X1 Society Int. GBO 31 Toring agriculture Control (1906) 1 April 2 Mary Home Sapiens CDNA clone IMAGE:1406935 3'	ACCHINES CONTROL OF LAST DITIONS CONFIDENCE	QV4-D LUOZI-SU IZBB-DI - DI OZI I GON SERGION	HYPOLIDE IOSE / Since the complete of since	MUS MUSCULOR CARDON CALIFORNIA OF WELL COMPANY LIVER CONTROL CALIFORNIA CONTROL CALIFORNIA CALIFORN	Z-NN10/3-ZZ0400-108-10/0 INVIOVS PROFILE CONTRACTOR OF THE PROFILE CON	Mus musculus microtubule-associated protein 2 (major), in the	601877026F1 NIH MGC_55 Home sapiens cutive cities invocative code 5	601877026F1 NIH MGC bb Homo sapiens curve cerre introduction and the product of the company of t	462/09.61 Soares_testis_NHT Homo sapiens cDNA clone IMAGE. 1334921 3 Sullinal IN BACTORE PROTEIN OF (HI MAN):	PINGEN FIX OF THIS SET THOM SERVENS CON Clone IMAGE:3866963 5	Comments mRNA for myomodulin neuropeotide precursor	Home seniens mRNA for KIAA0339 protein, partial cds	Compositions makes for KIAAR339 protein, partial cds	Figure September 10 13 genomic DNA 1166001-1485000 nt. position (6/7)	Pyrococcus Industrial Cologo and Population Population (Population Cologo and Population Population Cologo and Population Population Cologo and Population Population Cologo and Population Population Cologo and Population	Cea miss adult in Cost in the Art Home seniors cDNA clone IMAGE:3632190 5	CONSTRUCTOR IN TARCES A Homo sapiens CDNA clone IMAGE:2933740 3' similar to contains	L1.t1 L1 repetitive element;	historia Assares NFL_TGBC_S1 Homo sapiens cuina cione introck_5555 FC silling a communication of the communication	L1,t1 L1 repetitive element;	Homo sapiens chromosome 21 saginant no 210004	Organization will high Adv 10 As princes here Genomic 361 nt. segment 1 of 2]	Initiosatemic Involvity (1975 English English English English Francisco (RABEXS), mRNA	necessary Spares, pregnant uterus, NbHPU Homo sapiens cDNA clone IMAGE:1696982.3'	
Designation of the control of the co	Top Hit Database Source		T HUMAN			Т	Т	Т		ISSPROT	\neg	T HUMAN			EST HUMAN 60		Т	LIOMAIN					Т	EST HUMAN	EST_HUMAN L		EST HUMAN				TOTAL	1
5	Top Hit Acession No.	_	7.7					9	33.1		٦	7.1	8943	02 BF241924.1								-	-02 U62749.1	-02 BE394869.1	1.7E-02 AW573183.1		1.7E-02 AW573183.1	-02 AL163204.2	-02 AB004816.1	S74186.1	(N) 684/69/	1.7E-02 AI14/615.1
	Most Similar (Top) Hit BLAST E	1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.8E-02/	1.8E-02	1.8E-02 AW9363	1.8E-02		1.8E-02	1.8E-02		1.8E-02		1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.8E-02	1.7E-02	1.7E-02		1.7E-02	1.75	1.75			
	Expression Signal	4.1	0.83	1.32	1.73	1.71	0.94	0.39	1.17	5.02	69.0	16.0	0.8	0.49	0.49		2.41	1.51	1.37	2.31	2.31	1.59	3.32	1.86	2.17		2.17	3.41	10.5			1.4
	ORF SEQ ID NO:	25772				L			29764	32671		L	١_							36409	L	37535		26323	27230		1 27231			3 27752		2 28411
	Exen SEQ ID NO:	13137	13446	13892	14164	15394	15968	16810	17133		1	L	1	1	L.	L	22195	22611	22774	23181	l_	24211	24223	ı	44524		14524	1_	14837	1_		3 15762
	Probe SEQ ID NO:	336	970	1137	1416	2885	3205	4065	4396	6712	8020	8367	2440	2000	1000	2858	8542	8963	10128	11414	11414	11613	11626	88	702	3	1783	1864	2106	2291	2848	2896

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		\neg		т.					_	_	_			_	_	_	_		_	_								_		
Single Exon Probes Expressed in Brain	Top Hit Descriptor	hm45a04.x1 NCI_CGAP_RDF1 Homo sapiens cDNA clone IMAGE:3015534.3' similar to containe	MER/19.51 MER/19 repetitive element;	ac1904.s.1 Strategene ovary (#937217) Homo sapiens cDNA clone IMAGE:858927.3' similar to contain Air.	repetitive element; contains element MER24 repetitive element;	progress; 1 socials state inter splean 1NFLS Home saplens cDNA clone IMAGE:124847 6' gm08g07.x1 NC]_CGAP_Lu5 Home saplens cDNA clone IMAGE:1881276 3' similar to gb.X62359 ZINC	h/34e/3.x/ Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2933740 3' similar to contains	C. I. T. Tepetitive element;	wessenger KNA for anglerfish (Lophius americanus) somatostatin II	ovo reuz.si Soares testis_NHT Homo sapiens cDNA clone IMAGE:1640858 3'	Down Indregicus N-arginine dibasic convertase 1 (Nrd1), mRNA	molito septetts 939 to contig between AML1 and CBR1 on chromosome 21q22; segment 1/3 wg35f09.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo septens cDNA clone IMAGE:23g7113.3' similar to	contains Alu repetitive element;	Mossa Salres Tetal liver spleen INFLS S1 Homo saplens cDNA clone IMAGE:16726613	macaca lasciculars protein tyrosine phosphatase (PRL-1) mRNA, complete cds	Harin Septens neoulin (NEB), mRNA	Human apolipoprotein (a) gene, exon 1	Human apolipoprotein (a) gene, exon 1	Conserts Living	Ceenoningous eregans cCAF1 protein gene, complete cds	Homo ganiene service constituent (1.2) Homo septems cDNA clone DKFZp43410314 6	CMAL NINTORO ADVANCE OF UNINTERED (MISES), MRNA	Mycobacterium #the control of the companies and the control of the	myceacterian tuber culosis H3/KV complete genome; segment 13/182	Treponema maltophilum flab2, flab3 and fliD genes for flagellin subunit proteins and CAB and in the control of	LIVER CARBOXYLESTERASE 22 PRECURSOR (FOASSAN) VESTERASE SING CAT PROBATION COMPANY OF THE STATE OF THE COMPANY OF THE STATE OF THE COMPANY OF	LIVER CARBOXYLESTERASE 22 PRECIPSOR / FOASWAY (ESTERASE 22)	Homo saplens KVLOT1 gene	ne81406.s1 NCI CGAP Ewit Home conference ONA circumstance of the conference of the circumstance of the cir	Homo saplens mRNA for KIAA0634 protein, partial cds
igie Exon Pro	Top Hit Database Source	H	SWISSPROT		EST HIMAN	EST HUMAN	TO LIBERT	FIS TOWAN	FOT LINAN	TO LOUISIN	TA	TOT TO	Т	NOND P						T HUMAN		T HIMAN					SWISSPROT		EST HUMAN n	
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Page 163 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	Lassea sp. isolate IBd cytochrome oxidase III gene, partial cds; mitochondrial gene for mitochondrial product	IL3-CT0219-160200-063-C07 CT0219 Home Septens CDNA	Homo sapiens chromosome 21 segment nozi circi	Mus musculus major histocompatibility complex region NG2T, NG26, RY-226, INDO musususususususususususususususu KIFC1, Fas-binding protein, BING1, tapasin, RalGDS-like, KE2, BING4, bata 1,3-galactosyl transferase, and	RPS18 genes, complete cds; Sacm21 gene, partial>	QV2-PT0012-140100-030-f07 PT0012 Homo saplens cDNA	Mus musculus CD5 antgen (Cd5), mrwA	Candida albicans CaGCK3 gene, complete dus	Saccharomyces cerevisiae CADZ gene ior caumium resistance promin, compress cas	Saccharomyces cerevisiae CADZ gene for caomium teaskalice process, compress cas	Arabidopsis thaliana DNA chromosome 4, cong magment No. 20	Homo sapiens partial TUB gene for tubby (mouse) homolog and LMO1 gene for LIM domain only 1 protein	Human apoC-II gene for preproapolipoprotein C-II	Drosophila melanogaster enhancer of polycomb (E(Pc)) mRNA, complete cds	nf19g03.s1 NCI_CGAP_Pr1 Homo saplens cDNA clone IMAGE:914260 sImilar to SW:TELO_RABI1	P28294 TELOKIN. [1];	nf19g03.s1 NC_CGAP_PT1 Homo sapiens curva done innoces 1-1200 summa con 1-1200 p.29294 TELOKIN. [1];	Ggallus microsatellite DNA (LEi0260 (≍T16iiiE11))	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 20	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 20	qz98e10x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:2042442.5	Homo saplens transcription factor (HSA130894), mRNA	w27b07.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone invace: 245925 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 90	Homo saplens CACNA1F gene, exons 1 to 48	Homo sapiens CACNA1F gene, exons 1 to 48	MR4-TN0115-080900-201-012 INOT 3 India Sapiets COND.	HYPOLINE ICAL CALCOUNTBINGS OF ICAL CALCOUNT	Cyanoping a paracona of money Company (KIAA1009), mRNA	
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Page 164 of 536 Table 4

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	Top Hit Descriptor	ni11c04.s1 NCI_CGAP_Br2 Homo sapiens cDNA clone IMAGE:1029990 3' similar to contains Alu repetitive element;	Mycobacterium tuberculosis H37Rv complete genome; segment 88/162	Candida boldinii methanol oxidase (AOD1) gene, complete cas	Homo sapiens SPP2 gene for secreted phosphoprotein 44 preductor, excits 1-6	601078239F1 NIH MGC 12 Homo sapiens cunA cione IMAGE. 340424 I 3	Homo sapiens chromosome 21 segment no 2100 lo	Human IFNAK gene tor interteron alphanoeta receptor	Arabidopsis theliana P21J9.2 mKNA, complete cus	Homo sapiens sperm associated angen (Or Act), mixed	Homo sapiens chromosome 21 segment rio 210001	602129475F1 NIH MGC_56 Hamo sapiens clorik cione introci. 1200200 5	602128475F1 NIH MIGC 56 Home sapiens conve invade: "Zoozoo o	Mus musculus beta-sarcoglycan gene, complete cus	Human germline T-cell receptor beta chain TCRBV17S1A1T, TCRBV2S1, TCRBV10S1P, TCRBV29S1P, TCRBV19S1P, TCRBV16S1, TCRBV11S1A1T, HVB relic, TCRBV28S1P, TCRBV34S1, TCRBV14S1,	TCRBV3S1, TCRBV4S1A1T, TRY4, TRY5, TRY8, TRY7, TRY8, TCRBD1, TCRBU1S1, TCRBJ1S2,	Mus musculus chromosoma X contigB; X-linked lymphocyte regulated s gene, Zinc linger protein 27 v, zuro finger protein 92, mmxq28orf	Mins miscarifus chromosome X contigB: X-linked lymphocyte regulated 5 gene, Zinc finger protein 275, Zinc	finger protein 92, mmxq28orf	Homo sapiens basic transcription factor 2 p44 (bt/2p44) gene, partial cds, heuronel apoptosis intuolior) protein (naip) and survival motor neuron protein (smn) genes, complete cds	C.reinhardtii ribulose 1,5-bisphosphate carboxylase/oxygenase activase mKNA, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 46	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 46	ow06g05.x1 Soares_parethyroid_tumor_NbHPA Homo sapiens cDNA clone iMAGE:1648072 3' similar to	contains Alu repetitive element;	Homo sapiens human endogenous retrownus W gagles.37 / c gag (gag) gene, cumples cus	Mouse kidney androgen-regulated protein (KAP) gene, complete cus	Chlamydia trachomaus section of or or or or or or or or or or or or or	XXQ4e03.X1 Soares NPL GBC_S1 nomo saplens CDNA clone IMAGE:2815036 3	Worksup Names International Control of the Control
200	Top Hit Database Source	EST HUMAN	П			T HUMAN					П	EST_HUMAN	\neg	NT		TN	Þ		Ę	IN	N	LN	<u>LN</u>		EST_HUMAN	NT	L	N	EST HUMAN	EST_HUMAN
2	Top Hit Acesslon No.	1 4F-02 AA559030.1	AL022073.1	1.4E-02 M81702.1	2 AJ272265.1	2 BE544561.1	2 AL163218.2	2 X60459.1	1.4E-02 AF324985.1	11426968 NT				02 AF169288.1		02 U66061.1	A 040866 2	ALCHEOLO.	02 AL049866.2	02 U80017.1	02 M62962.1	AL161546.2	1.3E-02 AL161546.2		02 A1031593.1	02 AF156961.1	1.3E-02 M63707.1	1.3E-02 AE001304.1	1.3E-02 AW268563.1	-02 AW 268563.1
	Most Similar (Top) Hit BLAST E Value	1.4F-02./	1.4E-02/	1.4E-02	1.4E-02	1.4E-02	1.4E-02	1.4E-02	1.4E-02	1.4E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02		1.3E-02	20 10	20-36.1	1.3E-02	135	1.3E.	1.3E-02			1.3E-	1.3E			1.3E	1.3
	Expression Signal	5 47	1.94	0.75	0.84	2.27	0.76	2.2	1.37	2.36	2.21	2.23	2.23	1.31		0.93		10.7	1.34						4.9	1.65				3.97
	ORF SEQ ID NO:	og OC.		34634	<u> </u>			37267			27395					30223		30232	30533						33244			36040		36839
	Exon SEQ ID NO:	40004			L		23101		L_	24913	1_	١.		L	<u> </u>	17601	1	1/9/5	17075	1		1	١	L	20150		L	3 22826	L	3 23593
	Probe SEQ ID NO:	7000	8038	8796	9053	9300	10455	11985	12331	12601	1948	3208	3208	3954		4874		5166	2466	8	8405	2000	2000	8	7477	8380	10107	10178	10913	10913

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Single Exon Probes Expressed in Brain	Top Hit Database Source		Bacillus subtilis complete genome (section 14 of 21): from 2500451 to 2010000	Human herpesvirus 6B, complete genome	Homo sapiens V1b vasopressin receptor (VPR3) gene, complete cds	EST HUMAN 1.1 repetitive along 1024HK Home saplens oDNA clone IMAGE:381840 G similar to contains alonged.	1	Г	T	1	Т	T	Т	┑	Т		т	\neg	Human hereditary haemochromatosis region histone 24 III.	(HLA-H) gene, RoRet gene, and sodium phosphala transmission promised large hereditary haemochromatosis	7	HUMAN AV731704 HTF Hano septens a DNA clone HTFBHG11 g.	Rana rugosa mRNA for cairettculin, complete cds	Homo saplens wasart (WBSCR1) and wasars (WBSCR6) denes, complete and colors.	replication tactor C subunit 2 (RFC2) gene, complete cds	7	7	7	$\neg \vdash$		\exists	Home saplens fringe protein mRNA, partial ods	Inomo sapiens tringe protein mRNA partial cas
ngle Ex	Dat		2	Z	z	EST H	SWISSPROT	100	HOMAN	2 2	EST HUMAN	TOT	FOT HUMAN	ES HOMAN	ES HOMAN	EST HIL	FOT LINAN	3	į	2 !		-1	Z	Ė		N.		ESI HUMAN	HOMAN		SWISSPROT		
รัง	Top Hit Acess No.	13F-02 700117 4		1.3E-02 A E 162239 4	10000	1.2E-02 AA059299.1	-02 P38898	02 41483822 4	02 Al 163249 2	AV734704.4	1.2E-02 AW172350 1	T	T	T		1.2E-02 A 1668694.1	02 A 887378 4			1 2E-02 091328.1	ľ	T		AF045555.1	T	T	,	T	T		AE402642.4	AF193612.1 INT	
	Most Similar (Top) Hit BLAST E Value	13F-02	1.3E.02	135-02		1.2E-02	1.2E-02	1.2F-02	1.2E.02	1 2F.02	1.2E-02	1 2F.02	1.2E.02	1 2F-02		1.2E-02/	1.2E-02/		1 20 00	1 2 2 2 2	125.03	1 2F-02 726FB0 4		1.2E-02 A	1.2E-02 A	1.2E-02.H	1.2F.02 AV				1 2E.02 A	1.2E-02 AF	
	Expression Signal	4.	2.41	28.18		3.48	1.66	2.02	1.81	1.71	1.39	1.07	6.66	2.1		0.92	0.91		2.03					0.72	6.46		10.54			218			
	ORF SEQ ID NO:				00000	23/07	80007	26148	27637	27640	27901	27901		28691	-	28694	30045		30211		30358	31375		31767	32691	32952	32972	33222	_	33722	33917	33918	
	Exan SEQ ID NO:			25145	13446		3	13495	14904	14907	15163	15163	15863	16042	!	16045	1/409		17588	17704	17746	18461	_	18806	19645	19878	19897	20130		20591	20786	20786	
	Probe SEQ ID NO:	12337	12437	12607	345	8		721	2176	2178	2444	2842	888	3281	-	3284	2/2		4859	4981	5025	2666		_	-1	7192	- 1	7458		7896	8082		

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Top Hit Descriptor	yd72c08.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:113774 3'	Norwalk-like virus genogroup 2 gene for capsid protein, complete cds	Homo sapiens Spast gene for spastin protein	PERIOD CIRCADÍAN PROTEIN 1 (CIRCADIAN PACEMAKER PROTEIN RIGUI) (HPER)	C18119 Human placenta cDNA (TFujiwara) Homo sapiens cDNA clone GEN-557G06 5'	zm69e11.s1 Stratagene neurospithelium (#937231) Homo sapiens cDNA clone IMAGE:5309243'	H.sapiens LIPA gene, exon 4	H.saptens LIPA gene, exon 4	602018037F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4153808 5	za40e06.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295040 5'	tq95b10.x1 NCI_CGAP_Ov23 Hamo sapiens cDNA clane IMAGE:2216539 3' similar to SW:XPF_HUMAN Q92889 DNA-REPAIR PROTEIN COMPLEMENTING XP-F CELL ;	RC3-ST0197-120200-015-g11 ST0197 Homo sapiens cDNA	DKFZp686E0924_s1 586 (synonym: hute1) Homo sapiens cDNA clone DKFZp586E0924	Bacillus subtilis SpoVK (spoVK), YnbA (ynbA), YnbB (ynbB), GinR (ginR), glutamine synthetase (ginA),	Yna4 (yna4), YnaB (ynaB), YnaC (ynaC), YnaD (ynaD), YnaE (ynaE), YnaF (ynaF), YnaG (ynaG), YnaH (ynaH), YnaI (ynaI), YnaJ (ynaJ), xylan beta-1,4-xylos⊳	RC1-HT0256-100300-016-h07 HT0256 Hame saplens cDNA	QV3-BN0045-220300-128-h02 BN0045 Homo sapiens cDNA	C04803 Human heart cDNA (YNakamura) Homo sapiens cDNA clone 3NHC4040	NEUROGENIC LOCUS NOTCH 3 PROTEIN	zn24a01.r1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens cDNA clone IMAGE:548328 5	EST186494 Colon carcinoma (HCC) cell line II Homo sapiens cDNA 5' end	Homo septens T-box 5 (TBX5), mRNA	ab7711 s1 Stratagene fetal retina 837202 Homo sepiens cDNA clone IMAGE:853005 3' similar to contains Alu repetitive element	RC3-ST0197-120200-015-g11 ST0197 Hamo sapiens cDNA	MR3-CT0176-111099-003-e10 CT0176 Homo sapiens cDNA	CM2-HT0177-041099-017-h12 HT0177 Homo sepiens cDNA	oc22h08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1350495 3'	RC0-FN0025-250500-021-d02 FN0025 Homo sapiens cDNA	601649967R1 NIH_MGC_74 Hamo sapiens cDNA clone IMAGE:3933689 3'	HA0921 Human fetal liver cDNA library Homo sapiens cDNA
Top Hit Database Source	EST_HUMAN	TN	TN	SWISSPROT	EST_HUMAN	EST_HUMAN	IN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN		E	EST_HUMAN	EST HUMAN	EST_HUMAN	SWISSPROT	EST HUMAN	EST_HUMAN	N	NAMILIE TOT	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	T76987.1	AB031013.1	AJ246003.1	015534	C18119,1	AA070384.1	X75491.1	X75491.1	BF345263.1	N99523.1	A1653508.1	AW813796.1	AL048383.2		U66480.1	BE149611.1	AW996160.1	C04803.1	Q61982	AA082578.1	AA314865.1	11435505 NT	A A S S R 2 3 0 4	AW813796.1	AW846120.1	AW368128.1	AA806389.1	BE83556.1	BE968999.1	A1065086.1
Most Similar (Top) Hit BLAST E Value	1.2E-02		1.2E-02	1.2E-02	1.2E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02		1.1E-02		1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	4 1E-02			1.0E-02	1.0E-02	1.0E-02	1.0E-02	1.0E-02
Expression Signal	1.03	2.46	1.35	2.88	8.02	1.49	1.35	1.35	4.92	4.05	2.98	0.86	1.5		_	2.51	0,91	29.0	6,45	2.03	3.55	3.23	4.48	1,62	9.16	1.56	1.57	27	1.49	0.79
ORF SEQ ID NO:		35376				26662	27143	27144	27495		28924		30132		31799				34639	35682						26948		28494		29250
Exen SEQ ID NO:	21480	22192	2223	24559	l .	l	14444	14444	14766	15647	16269	16829			18837	20169	1		21492	22480	L	L	24484	1	<u>L</u> _	14260	15291			16611
Probe SEQ ID NO:	8788	9539	9570	12034	12615	1246	1701	1701	2031	2880	3513	4086	4778		6057	7497	8538	8721	8800	6286	9884	10900	44023	12678	Θ	1513	2577	3087	3257	3861

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Table 4

Page 169 of 536 Table 4 Single Exon Probes Expressed in Brain

Single Exon Probes Expressed in braili	Top Hit Descriptor	qh87c12x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1853974 3	Home saplens hypothetical protein r.L.1 1000 (1 L.1 1000), http://www.n.n.n.n.n.n.n.n.n.n.n.n.n.n.n.n.n.	UNITED STATES AND STATES STATES AND STATES OF	spliced spires approximate spiral spi	INTEGRIN BELAY, PRECOROON (INTEGRIN DELINA) (WESSELL AND AND AND OBEIN HESSELL AND AND AND OBEIN INSOR	COLLAGEIN ALT TA I(V) CIVILIA CONTROL	Rodatores Ft All Mig. 44 Homo sabiens cDNA clone IMAGE:3632181 5	Lower services preprogatatin (GAL1) gene, exchs 1, 2, and 3	Home soules propredatin (GAI 1) gene, excess 1, 2, and 3	Home Sapratis propriedate III (2012 / 3013) Society Cone IMAGE:3183161 3	DAY UTAKE 20120 011 AND HT0452 Home sepiens cDNA	FMI-HIVE COLOR FOR SAME AND SEALERS CDNA Clone IMAGE:413596 3' smilar to contains	Zh30e(3,31 Sodres prired yial 2,351 ii 3 153 days	Homo sapiens adenylosuccinate lyase gene, complete cas	Homo sapiens chromosome 21 segment 1521 was	Escherichia coli microcin 24 region, DNA binding protein (mabA), immunity protein (min, mino com 2.7 mino), and microcin transport protein (mtA, mtB) genes, complete cds	Homo sapiens SCL gene locus	HYPOTHETICAL 127 0 KD PROTEIN IN RAD24-BMH1 INTERGENIC REGION	HYPOTHETICAL 127 0 KD PROTEIN IN RAD24-BMH1 INTERGENIC REGION	CAAA_NN0119_300600-223-b05 NN0119 Homo saplens cDNA	ATTAMOSE PLACES Homo sapiens cDNA clone PLACE2000223 5	No. 170201 1. State Mistrompatibility complex region NG27, NG28, RPS28, NADH oxidoreductase, NG29,	KIFC1, Fas-binding protein, BING1, tapasin, RalGDS-like, KE2, BING4, beta 1,3-galactosyl transferase, and	RPS18 genes, complete cds; Sacm21 gene, partial>	Pyrococcus horikoshii OT3 genomic DNA, 287001-544000 nt. position (211)	PROBABLE PEPTIDASE Y4NA	Human BK virus (strain MM) genome. (Closely related to SV40.)	A californica (marine gastroped mollusc) neuropeptide gene (bag cell), exon 1, 5 end	Tursions fruncatus mRNA for p40-phox, complete cds	RASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN	\neg
le Exon Prot	Top Hit Database Source	EST HUMAN	L	EST HOMAN	Į.	SWISSPROT	SWISSPROT	IN L	EST HOMAN	Z	±N,	EST HUMAN	EST HOMAN	EST_HUMAN	NT	LN	FZ	FIN	TOGGGGWG	SWISSING TOUGGE	SWISSTROIL	TOWN HOLD	אלאוסט ומש		LN LN	NT	SWISSPROT	LN.	LN	Z LZ	1	SWISSPROT
Sing	Top Hit Acession No.	1242219.1	22570		9.0E-03 AF223391.1	26011	20908	18000.1	9.0E-03 BE395380.1	11144.1	11144.1	9.0E-03 BE348385.1	9.0E-03 BF351141.1	B.0E-03 AA723007.1	3 AF106656.1	8 0F-03 AL 163283.2	0 OF 02 1147048 4	440404	8.0E-03/AJ131010.1	-32044	P32644	03 BF363327.1	03 AU140261.1		AF110520.1	8.0E-03 AP000002.1	P55577	V01109 1	1447407.4		03 AB038207.1	.03 P98160
	Most Similar (Top) Hit BLAST E Value	9.0E-03 AI242219.1	9.0E-03	9.0E-03 A	9.0E-03	9.0E-03 P26011	9.0E-03 P20908	9.0E-03	9.0E-03	9.0E-03 L11144.1	9.0E-03	9.0E-03	9.0E-03	8.0E-03	8.0E-03	8 0E-03	100	0.00	١		8.0E	8.0	8.0E		8 OF-03	L	L		_	S.OF	8.0 F.O.	8.0E
	Expression Signal	0.57	8.0	1.05	0.65	0.47	1.44	2.07	1.57	1.55	1.55	2.37	23.46	4 06	36,32	20	3	0.83					1.09		2.82						2.03	0.63
	ORF SEQ ID NO:	33122	33132			35601	35620		36866							27847			ı				30420			30949	١		ı	32860		34619
	SEQ ID	20043	20051	20470	20841	22396	22413	23587	23615	24248	24248	25411	24983	l	┸	1				16418	16418	17089	17802	_		18235	1	1	_1	19795	20119	21473
	Probe SEQ ID NO:	7367	7371	7774	8447	9745	9762	10907	10935	11651	11651	12411	12703	9	890	300	\$ 7	2960	3353	3865	3665	4350	5083		-	5436	010	9851	9820	7107	7442	8781

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	-				เริ่	ngle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E	Top Hit Acession		Top HIt Describer
	J			Value		Source	
8808			3.29	8.0E-03	8.0E-03 AW808602 1	FOT LINAANI	MP4 ST0444 44450
8816	3 21508		0.49	8.0E-03	8.0E-03 At 139075 2	TWO LEN	MRT-S10111-111199-011-h06 ST0111 Homo seplens cDNA
8878		34713	0.58	8.0E-03	789956 NT	Į.	Min minoring 6 in 10 CT CT 1168 complete genome; segment 2/8
9848	22498		4.63	8.0F-03	REDARKO	COT UNIVERSE	was musculus fusion 2 (human) (Fus2), mRNA
10686	23357	36597	1.36	8.0E-03	8.0E-03 BF788441 4	EST HUMAN	CV1-5 10677-040400-131-903 BT0677 Homo saplens cDNA
10906	23586		3.58	8.0F-03	03 749652 1	NT TOWAIN	OU 147 001 BFT NIH MGC 68 Homo sapiens cDNA clone IMAGE:3878405 6
11715	24309	37632	4.74	ROF-03	O3 AEABAGBO 4	L L	S.cerevisiae chromosome X reading frame ORF YJR152w
11814	24402		22.71	8.0F-03	l	For Links	Inomo sapiens melanoma-associated antigen (MAGE-C1) gene, complete cds
11853	24437	37779	1.36	8.0F-03		EOT LINAN	2832911.r1 Soares retina N2b4HR Homo saplens cDNA clone IMAGE:360716 5:
11933	24491	-	174	8 OF 03		ES : TOWAIN	OVZU13841F1 NCI_CGAP_Bm64 Homo saplens cDNA clone IMAGE:4149418 5
	L				, conson		Uryciolagus cuniculus elF-2a khase mRNA, complete cds Homo santans & BCG1
11880			1.74	8.0E-03	_	N T	complete ods
8/8	.1	26098	16.15	7.0E-03	7.0E-03 AF097183.1	L	Cholosophilin penum HC 40 sees
978		26097	16.15	7.0E-03/		Ł	Chaptersocial and the value of the complete cas
828		26387	3.67	7.0E-03/		Į	Official move of the first of the complete cds
1092	13852	26511	3.48	7.0E-03	7.0E-03 AV7317121	FOT LIMAN	Aviation in the guaranter as GS 21 mRNA, partial ods
-	-					NUMBER	FORKHEAD BOX PROTEIN DS (LINES ICH TEXASOCIATED CO.
245	14091		2.67	7.0E-03 Q61060		SWISSPROT	NICE FAR FACTOR & EDELLEAD LONG LINES FOLLOW PACTOR GENESIS) (HEPATOCYTE
1374	14122	26797	6.71	7.0E-03	7.0E-03 AA668298.1	EST HUMAN	ab79h0a s1 Shatagana fotal and control 2) (HFH-2)
1491	14238	26924	3.37	7.0E-03		FST HIMAN	WORLD OF SECURITY OF STATE OF STATE OF SECURITY OF STATE
1735	14477	27175	1.24	7.0E-03	T	EST HUMAN	FST38983 MACE: 1 GBC: S1 Homo saplens cDNA clone IMAGE:2813739 3
1735	14477	27176	1.24	7.0E-03 A		Т	ESTRACACE MACE resequences, MAGA Homo saplens cDNA
2254	15589	27722	1.86	7.0E-03 P	l	SWISSPROT	HIGHDINE BIOL O'S CORDOTTE THE FEET OF SEPTEMBERS CONA
3548	16301	28951	0.71	7.0E-03 A		Т	034h0 vi sprenteri nitelia precionali in pre
3749	16502	29137	0.8	7.0E-03 A	7.0E-03 AW44463.1	T	UI-H-BI3-ekb-o-10-0-UI:s1 NCI CGAP Sub5 Home senien: CDNA class (NA OF STREET)
3782	16544	29179	132	7 00 00 1	A 1740000 444		S LADSS / Z JACKINI BIDIO CATO STOCKS
4000	16502	29137	0.83	7 SO TO TO	T.	7	Rattus norvegicus neuronal nicotinic acetylcholine receptor subunit (Alpha10) mRNA, complete cda
4560	17295		124	7.0E-03.A		ICMAN	U-H-Bi3-akb-c-10-0-Ui,s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2733891 3'
4929	17657		212	7 05 03 7	T	HOMAN	nnsdatus y1 NC_CGAP_GU1 Homo septens cDNA clone IMAGE:2969936 6
					1	Z	Homo saplens chromosome 21 segment HS21C078
6229	18521		0.75	7.0E-03/H	H741084 · E	EST HUMAN	y/82g01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:211824 6' similar to
6021	25083		4.9	7.0E-03 A	ļ	T	BCA CERSES SEALES
6222	18996	31972	1.47	7.0E-03 W	T	Т	7472410 15 Sec-20-4W-U18-008 CT 0286 Home sapiens cDNA
					1	7	2033110.11 Soares, Tetal heart NPHH19W Homo sepiens cDNA clone IMAGE:342475 5'

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Table 4

Exan SEQ ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:	ORF SEQ Expression (Top) Hit Top Hit Acession Database In Brain Top Hit Descriptor Top Hit Descriptor	Value	32207 3.44 7.0E-03 AA327129.1 EST_HUMAN EST30674 Colon I Homo sepiens cDNA 5 end	32237 0.75 7.0E-03 BE857385.1 EST HUMAN Q13387 HYPOTHETICAL PROTEIN 98/10-3 3. Similar to TR:Q13387	1.67 7.0E-03 BE928133.1 EST HUMAN	5.48 7.0E-03 Z35838.1 NT	5.48 7.0E-03 Z35838.1 NT	2.47 7.0E-03 BE175667.1 EST HUMAN	7.0E-03 AF281074.1 NT		0.72 7.0E-03 N52378.1 EST HUMAN	2.57 7.0E-03 P48982 SWISSPROT	2.57 7.0E-03 P48982 SWISSPROT	7.0E-03 AV687379.1 EST HUMAN	0.77 7.0E-03 AI799734.1 EST HUMAN	2.63 7.0E-03 AB008852.1 NT	1.71 7.0E-03 AJ004862.1 NT	7.0E-03 AJ004862.1 NT		EST HUMAN	7.0E-03 Y17455.1 NT	1.72 7.0E-03 AL163300.2 NT Homo sapiens chromosome 21 segment HS21C:100	26637 12.34 6.0E-03 AW 511148.1 EST_HUMAN SW.PXR_HUMAN O75499 ORPHAN NICLEAR RECEPTOR DXD.	26638 12.34 6.0E-03 AW511148 1 FST HIMAM SW-PYD HIMAM OTF AS OF DELAY NO. 12.34 6.0E-03 AW511148 1 FST HIMAM SW-PYD HIMAM OTF AS OF DELAY NO. 12.34	1.3 6.0E-03 AF112374.1 NT	3.36 6.0E-03 AA759135.1 FST HIMAN	3.36 6.0E-03/A4759135.1 FST HIMAN	6.0E-03 H75690.1	1.31 6.0E-03 AF190338.1 NT Notoncus sp. evtochrome c oxideses enhants it save avaitable at a contraction of the contraction of	
Exon SEQ ID OR SEQ ID NO: 19211 19211 19237 19504 20097 20097 20097 2273 22273 22273 22273 22273 22273 22273 22273 222657 23657 23657 23657 24890 2540	<u> </u>									0.7				1.3.					1.2	1.75	1.81	1.72						2.22	1.31	
			┙			Ц		П		2				25		\bot			12	2	g	او						2	0	
		- 1					L.I							_(. 1	- 1	- 1	- 1		_1		1		1					

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. · · · · · · · · · · · · · · · · · · ·	Top Hit Descriptor	Fugu rubripes zinc finger protein, isotocin, fatty acid binding protein, seplapterin reductase and vasatocin genes. complete cds.	20/3/11/1 Soares Darathyroid Himor NAHBA Homo somitone 20NA size 11/407-2003-201-	UI-H-BI4-apm-c-06-0-UI-81 NCI COAP Subs Home content about Less INA Content of the Content of th	-a07 R Todor Home serious ability	Section of the september of the section of the september	-b10 CT0204 Home saniens CDNA	15 Home sations cDNA close IMAGE 2050512 El	NHT Home series con class MAGE 462013 6	EST27116 Cerebellum Homo septiens CDNA 6' and circles to EST	EX PROTEIN 4 (SOD 4 DEOTERN)	control (OCT) TROLEIN)		18 Home seriese CONA Almo IMAGE 2020430 EI	Homo septems CDNA A' and	Homo ceniano chilla Al and	Homo saplans felomerase reasons francociotinos (TEDT) esse	PARSO danso plass (1 EM) gene, exons /-10 and complete cds	ow13a04.x1 Soares_parathyroid_tumo_NbHPA Home expiens cDNA clone IMAGE:1646670 3' similar to	and I Money desirent,	36 Homo saniene cDNA close IMA OE 30 E00 20 E1	oenhelitis (SSPE) virus mRNA for frision protein	122602.x1 NCI_CGAP_KId11 Homo sapiens cDNA clone IMAGE:2131202 3' similar to SW:R13A_HUMAN P40429 60S RIBOSOMAL PROTEIN 13A		Homo septens okedato acid-Inducible and cAMP-regulated phosphoprotein 19 (ARPP-19) mRNA, complete	ate placemid hEV/1 DNIA	Shearing privile Company of the Comp	zince finger probable E. (14011 (F. (14011) - DNA	te91c12.x1 NCI_CGAP_Pr28 Home sepiens cDNA clone IMAGE:2094070 3' similar to TR:000519 000519 FATTY ACID AMIDE HYDROLASE.:
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Fugu rubripes zinc finger protein, isotocin, fatty acid binding protein, sepie genes. complete cds	1 Sogres parathyroid filmor NISHBA Homo somions 20NIA - 1-1-	apm-c-06-0-Ulls 1 NCI CGAP Subs Home septems abus all	RC1-BT0606-260400-014-a07 RT0608 Domo soulong ablita	Mus musculus diucosamina-8-phosphate deaminase (Good) michila	RC0-CT0204-240999-021-510 CT0204 Homo saniens CDNA	600942904F1 NIH MGC 15 Homo saniens CDNA close IMAGE: 2050543 F1	0433c11.x1 Soares feetla NHT Homo capiens chald class 1440 E1462013.3	3 Cerebellum II Homo seniens cDNA 5' and similar to EST	SYNAPTONEMAL COMPLEX PROTEIN 4 (CCD-4 DEOTEIN)	Varida virus, complete genome		601112353F1 NIH MGC 18 Homo seniens cDNA class (MA DE 2252412 E)	EST 11949 Uterus tumor I Homo senions CDNA A' and	EST 1949 Utens timor I Homo seniors CDNA 8' and	ens telomerase reverse transcriptons (TEDT)	RAS-RELATED PROTEIN RAD-20	ow13a04.x1 Soeres, parathyroid, tumor_NbHPA Homo eaplens cDNA clor	RCO-UM0051-210300-032-02 UM0054 Home Control Control	801454915F1 NIH MGC 66 Homo samens china class INV GE 19886578 E	Subacute sclerosing panencephalitis (SSPE) virus mRNA for fusion profein	122c02.x1 NCI_CGAP_Kld11 Homo sapiens cDNA clone IMAGE:2131202 P40429 60S RIBOSOMAI PROTEIN 1326	Bacillus subtilis fenD gene	ens okadalc acid-inducible and cAMP-regulated phosphoprotein	M.thermoformicicum complete plasmid nEV/1 DNA	EST374237 MAGE reseduences MAGG Home series CONIA	Homo saplens hypothetical zinc finder profile El 14011 (Fl 14011) mBNA	TEATTY ACID AMIDE HYDROLASE:
robes Expr		Fugu rubri	Т	Т	Т	Т	Т	Г	T	Т	Т	Variola viru	T	Г	Т	Т	Т	RAS-RELA	ow13a04.x	RCOLIMO	801454915	Subacute s	#22c02.x1	Bacillus sul	Homo saple	M.thermofo	EST374237	Homo saple	teg1c12x1 FATTY ACI
gle Exon Pr	Top Hit Database Source	FZ	EST HUMAN	EST HUMAN	EST HUMAN	LZ	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	LN	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	N IN	SWISSPROT	HEMAN	EST HUMAN	EST HUMAN	NT	EST HUMAN	NT	L	Ł	EST HUMAN	1	EST_HUMAN
Sin	Top Hit Acession No.	03 U90880.1		-		54029	03 AW847284.1	03 BE250108.1	03 A1016833.1	L		6.0E-03 9627521 NT		93.1	Ī		Γ	6.0E-03 P17964		6.0E-03 AW799337.1		6.0E-03 D10548.1				Γ	1.4	5814	AI420786.1 E
	Most Similar (Top) Hit BLAST E Value	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03,	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03 /
	Expression Signal	1.18	1.19	3.68	1.08	1.22	0.76	1.29	1.84	5.67	2.68	0.67	1.16	0.57	0.61	0.61	0.69	0.62	6.9	2.17	1.58	7:26	213	0.86	1.14	0.59	1.75	2.54	1.26
	ORF SEQ ID NO:	28794			29079	29147	29294			30013	30407	31802	32676	30559	32905	32906	33318	33479	33578	33695		35176	-	35928		36166	36575		36680
	Exon SEQ ID NO:		16290					ı	1	17381	17792	25084	19633	17924	19836	19836	20216	20365	20453	20569	20640	22004	22591	22710	22842	22951	23336	23402	23435
	Probe SEQ ID NO:	3377	3534	3652	3682	3759	3902	3838	4331	4647	5073	6061	6718	6755	7149	7149	7646	7702	7977	7874	7945	9464	9943	10062	10194	10304	10645	10713	10760

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Single Exon Probes Expressed in brain	SEQ ID NO: Signal No. Signal Value No. Source	23435 36881 1.26 6.0E-03 AI420786.1 EST_HUMAN	23583 4.6 6.0E-03 U14556.1 NT	23584 36833 2.81 6.0E-03 BE737895.1 EST_HUMAN	24563 3.25 6.0E-03 AF010496.1 INT	25474 6 69 6.0E-03 AE000833.1 NT	25235 3.17 6.0E-03 U30790.1 NT	24898 1.64 6.0E-03 BE788019.1 EST HUMAN	24910 1.68 6.0E-03 AJ245480.1 NT	13432 26072 2.7 5.0E-03 L25105.1 NT	13432 26073 2.7 5.0E-03 [25105.1 NT	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	13432 20072 3.73 3.02=30 2.02=3	ĻΝ	1.15 5.0E-03 AJ010457.1 NT	28135 2.5 5.0E-03 AB033006.1 NT	28351 0.88 5.0E-03 BE266057.1 EST_HUMAN	15898 28543 3.82 5.0E-03 T87623.1 EST_HUMAN	15915 2.83 5.0E-03 AL161491.2 NT	15927 28575 1.3 5.0E-03 R71794.1 EST_HUMAN	16033 1.12 5.0E-03 AJ297357.1 NT	16440 29082 4.03 5.0E-03 AF147449.2 NT	16494 29129 0.85 5.0E-03 U38914,1 NT	16704 EST_HUMAN	16494 29129 0.82 5.0E-03 U38914.1 NT	17304 29931 0.73 5.0E-03 AJ131016.1 NT	17404 30039 1.17 5.0E-03 A1752367.1 EST_HUMAN	TORIGE SAMES ROE OF PARENT SWISSPROT	Probe SEQ ID 8 NO: NO: 10750 10903 10903 10904 12042 12588 12588 12588 155 2588 2588 2588 2	Exon SEC ID NO: 23486 23583 23583 23583 23584 24683 24683 24898 248910 13432 13432 13432 13848 15502 15505 16649 17004 17004 17004 17004		Signal Signal 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	Most 25 B S S S S S S S S S S S S S S S S S S		Top Hit Database Source Source Source Source NT NT EST_HUMAN NT NT NT NT NT EST_HUMAN SWESPROT	Top Hit Descriptor Top Hit Descriptor FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FATTY ACID AMIDE HYDROLASE: FAULTY ACID AMIDE FOR HYDROLASE ACID AMIDE HYDROLASE ACID AMIDE ACID AMI
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Page 174 of 536 Table 4 Single Exon Probes Expressed is

	IOLESTERASE NG ENZYME FAF- E 9, Y					EST	
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| Top Hit Descriptor | PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE FAF-Y (UBIQUITIN THIOLESTERASE FAF-Y) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE FAF-Y) (DEUBIQUITINATING ENZYME FAF-Y) (ABIQUITINATING ENZYME FAF-Y) (ABIQUITIN-SPECIFIC PROTEASE 9, YOHROMOSONAE) | Chicamodowich Chicamodow & Bao Confirm of Confirm Chicamodow & Bao Confirm of Confirm Chicamodow & Confirm Chica | 600044584T NILL MCC 47 December 1 | Mus miscuilis AMP1 ages for 8 characteristics | Turslops fruncatus mRNA for bathanker complete cds | EST03012 Fetal brain, Stratagene (cat#936206) Homo saplens cDNA clone HFBCR93 similar to EST | RC3-CT0265-031089-014-07 CT0266 Umm comittee on the | Home series MASI 4 - Bits
 | RCB CT0281-081190-014 A A C OT0281 U | Poe Citosa Addes 611-Add CIO281 Formo sapiens con A | RCC-C 10261-081198-011-A05 C10281 Homo saplens cDNA | Maries complement state (ABS) - Bit A | Echanolis cal conceptor (CRZ) mRNA, 3 and | Raphit iteradokin (1101) was own 4 | SOFI PROTEIN
 | Plasmodium barahai 58 kDa ahasahamatsia mBNA and and and and and and and and and and | ROSTOSTOSOSOSOSOS STOSTO PER PER PER PER PER PER PER PER PER PER | ni46h10 s1 NCI CGAB Pro Home samine s | Homo sepiens PRO0471 protein (PRO0474) mBNA | ag49c10.81 Gessler Wilhis fumor Home senione CDNA close 1446 CE 14176560 21
 | 694F Heart Homo saplens cDNA clone 694
 | хл59g05.x1 Scares_NHCeC_cer/ical_tumor Homo saplens cDNA clone IMAGE:2898040 3' similar to
confains L1.12.L1 mpetitive element | xn69g05.x1 Soeres_NHCeC_central_tumor Homo saplens cDNA clone IMAGE:2698040 3' similar to | Wh09e04 rd Strategeon planning (#1007000) | Mus musculus handbattral and a MMCs. 4267 (100 pp. 100) | 248c04.7 NCI CGAP Bridg Home content of the city of content of | Mus musculus genomic fragment 279 Kh. chomoses 7
 | Mus musculus genomic fragment, 279 Kb, chromosome 7 |
| Top Hit
Database
Source | SWISSPROT | LN L | EST HUMAN | | Ę | FST HIMAN | EST HUMAN | I-N
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| Top Hit Acession
No. | 000507 | AE002234.2 | BE300091.1 | AB025024.1 | AB038267.1 | T05124.1 | 4W854327.1 | AB016816 1
 | AW855907.1 | W855907 1 | 48982 | AB1132.1 | 90723.1 | |
 | | l | Γ | 2557 |
 |
 | | | | 0946753 | |
 | П |
| Most Similar
(Top) Hit
BLAST E
Value | 6.0E-03 | 5.0E-03 | 5.0E-03 | 6.0E-03 | 6.0E-03 | 5.0E-03 | 5.0E-03 | 6.0E-03
 | 6.0E-03 | 5 OF -03 / | 5.0E-03 | 5.0E-03 | 5.0E-03 | 5.0E-03 | 5.0E-03
 | 5.0E-03 L | 5.0E-03 A | 6.0E-03 | 5.0E-03 | 5.0E-03
 | 5.0E-03 T
 | 5.0E-03 | 6.0E-03 A | 5.0E-03 | 5.0E-03 | 5.0E-03 B | 5.0E-03 A
 | 5.0E-03 A. |
| Expression
Signal | 2.97 | 0.91 | 7.56 | 7.45 | 0.64 | 0.73 | 1.21 | 7.5
 | 0.48 | 0.48 | 3.29 | 6.63 | 10,1 | 0.71 | 0.45
 | 0.89 | 0.7 | 0.45 | 0.51 | 0.48
 | 4.99
 | 3.42 | 3.42 | 1.89 | 1.47 | 3.54 | 1.53
 | 1.53 |
| ORF SEQ
ID NO: | 31694 | | | 30578 | | 33143 | | 33442
 | 33948 | 33949 | 33965 | | 34545 | 34676 | 35315
 | 36596 | 35716 | 35923 | 36091 | •
 | 1
 | 36785 | 36786 | 36900 | 36956 | | 37697
 | 37698 |
| | | 18769 | i | | | 20065 | 20170 | 20331
 | 20813 | 20813 | 20831 | 21201 | 21398 | 21530 | 22135
 | 22390 | 22621 | 22705 | 22879 | 23023
 | 23314
 | 23539 | 23539 | 23647 | 23693 | 23982 | 24365
 | 24365 |
| Probe
SEQ ID
NO: | 6963 | 6988 | 6489 | 6746 | 6940 | 7385 | 7498 | 7867
 | 8119 | 8119 | 8137 | 8209 | 8706 | 8838 | 9482
 | 9739 | 9871 | 10057 | 10231 | 10377
 | 10621
 | 10859 | 10859 | 10971 | 11021 | 11303 | 11774
 | 11774 |
| | Exon ORF SEQ Expression (Top) Hit Top Hit Acesston NO: Signal BLASTE No. Source | Exon ORF SEQ Expression (Top) Hit Top Hit Acession Database NO: Signal BLASTE No. Signal Value Source Value Source 18735 31694 2.97 6.0E-03 000607 SWISSDROT | Exon ORF SEQ Expression (Top) Hit Acession Top Hit | Exon No.: ORF SEQ ID NO: Expression Signal Top Hit Acession (Top) Hit Top Hit Acession Signal Top Hit Acession Database No: Top Hit Acession Source Source Source Source Source Source Source Nalue 18735 31694 2.97 6.0E-03 O00507 SWISSPROT SWISSPROT SOURCE NT EST HIMAN IN IMAN IMA | Exon No.: ORF SEQ ID NO: Expression Signal Top Hit Acession (Top) Hit Top Hit Acession Signal Top Hit Acession Database No: Signal Top Hit Acession Signal Top | Exon
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SWISSPROT 18769 0.91 6.0E-03 AE002234.2 NT 17816 7.56 6.0E-03 AB0250224.1 NT 19422 0.64 6.0E-03 AB0325024.1 NT | Exon No: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Acesston No: Top Hit Acesston Source No: Top Hit Acesston No: Top Hit Acess | Exon No: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Acesston No: Top Hit Acesston No: | Exon No.: ORF SEQ ID No.: Expression Signal (Top) Hit Top Hit Acession No. Top Hit Acession Source National No. Top Hit Acession Source National No. Top Hit Acession Source National Na | Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acesston Source Top Hit Acesston Source Top Hit Acesston Source NO: Signal BLASTE No. No. Source Source 18735 31694 2.87 6.0E-03 O0507 SWISSPROT 18789 0.91 6.0E-03 AE002234.2 NT 17916 30576 7.56 6.0E-03 AB025024.1 NT 20065 33143 0.73 6.0E-03 AB038267.1 NT 20370 1.21 5.0E-03 AB058027.1 EST HUMAN 20170 1.21 5.0E-03 AB058027.1 EST HUMAN 20371 33442 7.5 6.0E-03 AB058027.1 EST HUMAN 20371 33442 7.5 6.0E-03 AB058027.1 EST HUMAN 20371 33442 7.5 6.0E-03 AB058027.1 EST HUMAN 2081 33148 0.48 6.0E-03 AB051816.1 NT | Exon NO: ORF SEQ ID NO: Expression Signal Top Hit Acession No: Top Hit Acession Source National Signal Top Hit Acession Source National Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession Source National No: Top Hit Acession No: | Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acession Patabase Top Hit Acession Source Top Hit Acession Source NO: 1D NO: Signal BLASTE No: No: Source Source 18735 31694 2.97 6.0E-03 O0507 SWISSPROT 18789 0.91 6.0E-03 AE002234.2 NT 19244 7.56 6.0E-03 AE002034.1 NT 19422 7.45 6.0E-03 AB025024.1 NT 20065 33143 0.73 6.0E-03 AW056327.1 NT 20170 1.21 6.0E-03 AW0564327.1 EST HUMAN NT 20331 33442 7.5 6.0E-03 AW0564327.1 EST HUMAN 20813 33949 0.48 6.0E-03 AW0565007.1 EST HUMAN 20813 33849 0.48 6.0E-03 AW0565007.1 EST HUMAN 20831 33849 0.48 6.0E-03 AW0565007.1 EST HUMAN 20831 33866 3.29 | Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acession Patabase Top Hit Acession Source Top Hit Acession Source NO: 10 NO: Signal BLASTE NO: NO: Source Source 18735 31694 2.97 6.0E-03 O00507 SWISSPROT 18789 0.91 6.0E-03 AE002234.2 NT 17815 30578 7.45 6.0E-03 AE002234.2 NT 17816 7.45 6.0E-03 AE002234.1 NT 19422 7.45 6.0E-03 AE002234.1 NT 20065 33143 0.73 6.0E-03 AE025024.1 NT 20170 1.21 6.0E-03 AW864327.1 EST HUMAN 20331 33442 7.5 6.0E-03 AW865907.1 EST HUMAN 20313 33849 0.48 6.0E-03 AW865907.1 EST HUMAN 20831 33849 0.48 6.0E-03 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3.3443 0.48 6.0E-03 AW865907.1 EST HUMAN 21398 34645 0.48 6.0E-03 AW865907.1 NT 21398 34676 0.71 6.0E-03 AW86900.1 NT 22139 36956 | Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acesston Source Top Hit Acesston Top Hit Top Hit Acesston Source Top Hit Acesston Value Top Hit Top Hit Acesston Source 18735 31694 2.97 6.0E-03 AE002234.2 NT Source Source 18769 0.81 6.0E-03 AE002234.2 NT NT NT 19224 7.56 6.0E-03 AE002234.1 NT NT 20065 33143 0.73 6.0E-03 AE002234.1 NT 20170 1.21 6.0E-03 AE002234.1 NT 20170 0.48 6.0E-03 AE038267.1 EST HUMAN 20170 1.21 6.0E-03 AW865907.1 EST HUMAN 20170 33442 7.5 6.0E-03 AW865907.1 EST HUMAN 21081 33545 0.48 6.0E-03 AW865907.1 EST HUMAN 21082 3366 0.48 6.0E-03 AW865907.1 NT 21080 3366 0.48 6.0E-03 AW865907.1 NT 21396 0.46 0.0E-03 AW865907.1 NT 22139 <td< td=""><td>Exon ORF SEQ Expression Signal (Top) Hit Top Hit Acession Top Hit Top Hit Acession Source Top Hit Top Hit Acession Top Hit Top Hit Acession Source Top Hit Top Hit Acession Source 18735 31694 2.97 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Single Exon Probes Expressed in Diam	Top Hit Descriptor	Gailus gailus glyceraidehyde-3-phosphate dehydrogenase minnin, completa cos	Brugia malay Y chromosome marka	Human pro-alpria 1 type in College (Cont.) 3 - College (MAGE:809548 3' similar to	SW.DXX. MOUSE 194688 TO PROPERTY OF THE PROPER	6020/7/174F1 Nin Midd of Halifo Substitution	COLLAGEN ALPHA 1/VII) CHAIN PRECURSOR (LONG-CHAIN COLLAGEN) (LC COLLAGEN)	11.HE.RND-akc-h-04-0-UI.1 NIH MGC 50 Homo sapiens cDNA clone IMAGE:3076831 5'	une 1904 s.1 Spares infant brain 1NIB Homo septens cDNA clone IMAGE:35988 3'	283 HOTH Spares NFL T GBC S1 Homo sapiens CDNA clone IMAGE: 1582568 3'	CITY STATES TO STATES THE HAND SERVICES CON CITY CONTROL TO STATES THE PARTY OF THE	ygo red-si sogles interior and BT0333 Home sapiens cDNA	RACA-B 1035-11 1010-12 - From Septimes CDNA clone IMAGE:510998 57	201auo. 11 Juna 4 470 Ann 022 Christian September CDNA	KCG-UMUU 14-17 U4-04-56-56-1 Umo saniens cDNA clone IMAGE: 701736 5	2538801.11 NOT COAT GOOD TONG CARE ADCAKEOG 5	TAY COOK OF THE CONTRACT OF TH	Rattus novegious type 1 astrocyte and offactory-limbic associated protein AT1-46 mRNA, complete cds	201avo.r.1 Suraisgere court, 100 227/1000 CDNA clone IMAGE:3638510 5	SUISO41017 LINE CONTRACTOR INCOME SECTION SECTIONS SECTIO	No construction of the second	Homo sapera Azo regul near Azo comming protein kinase I (CAMKI), creatine transporter (CRTR), protein LT48 (PRLT8), and construction dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM protein (CDM), adrenoleukodystrophy protein > CDM protein (CDM), adrenoleukodystrophy protein > CDM	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal	protein L18a (RPL18a), Ce2+/Calmodulin-dependent protein kinase I (CAMKI), creame transporter (Chin), CDM), adrenoleukodystrophy protein >	Lome engine notwintermine-containing C14ORF4 gene	Livra series polyclutamine containing C14ORF4 gene	Home same chromosome 21 segment HS21C084	Date: H70240-151239-003-h08 H70340 Homo saplens cDNA	PM1-HT0340-151299-003-h08 HT0340 Homo sapiens cDNA	X198104.x1 NCI_CGAP_Co18 Homo saplens cDNA clone IMAGE:2665279 3'	
e Exon Probe	Top Hit Database Source			Ż		\neg	1	T	Т	Т	Т	Т	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	LZ	EST HOMAN	EST HUMAN	EST HOMAN	Ŀ		<u>.</u>	12	Z	Ž	INVIEW FOL	FOT LIMAN	EST HIMAN	E01 110111
Sing	Top Hit Acession No.	\prod				1	99.1	202388	4.0E-03 AW500196.1	03 R46482.1	4.0E-03 AA939339.1	03 R46482.1	03 AW749101.1		_{	1	03 AV708305.1	03 U33472.1	03 AA099777.1	03 BE410556.1	-03 AW 794740.1	6 44	03 035111.4		4.0E-03 U52111.2	4.0E-03 AJ277385.1	-03 AJ277365.1	4.0E-03 AL163284.2	4.0E-03/BE154134.1	4.0E-03 BE13413411	AW 180420.1
	Most Similar (Top) Hit BLAST E Value	5.0E-03	5.0E-03 /	5.0E-03 L10347.1	5.0E-03	5.0E-03	5.0E-03	5.0E-03 Q0238B	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	L	4.UE-U		4.0E-03	4.0E-03	4.0E-03				
	Expression Signal	9.26	4.11	2.52	1.67	2.78	4.2	1.76	2.98	2.29	2.69	2.03	4.64	29.46	2.4	1.57	1.52	2.23	10.56	2.49	1.64		1.97		1.97		3				0.97
	ORF SEQ ID NO:	1					30980		25675		25997	26298		26544		<u> </u>		27178			27742		78030		3 28031	5 28140				2 28635	
	SEO D	25367	1	24792	24811	25183	24940	L.	1_	13117	13369	13626	_	L	1_	L	14327	14479	L		L	ŀ	15293		15293	3 15405	5 15405	L	15982		1 16277
	Probe SEQ ID NO:	42478	12307	12409	12441	12467	12643	12662	228	313	88	857	ğ	1128	1148	1280	1581	1737	8	2244	2276		2579		2579	2696	2696	2701	3219	3219	3521

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Single Exon Probes Expressed in Brain	1ft ase Top Hit Descriptor	Т	_[.	Т	Mus musculus fumor susceptibility protein 101 (tsg101) gene, complete cds	Homo saplens TNNT1 gene, exons 1-11 (and jained CDS)	AN L1 repetitive element:	T	Т	Rath s procedure both sets along 1 (anong 17) mRNA, complete cds	Transa revegues beta-carenin binding protein mRNA, complete eds	Т	7	T	7	Т	T	T	Lycopersion esculentum knotted 3 protein (TKn3) mRNA, complete eds	nomo sapiens chromosome 21 segment HS21C078	T.	т	Т	H.sapiens hoolX cane	П	7	Unictyostellum discoldeum AX4 development protein DG1122 (DG1122) gene, partial ods		7	Homo saplens chromosome 21 segment HS21C009	П	7	Arabidopsis thallana DNA chromosome 4, contig fragment No. 55	Trumo suprens chromosome 21 segment HS21C081
igle Exon	Top Hít Database Source	100	EST HOMAN	SWISSPRO	Z	2	EST HUMAN	EST HUMAN	N	Į	TOGGSSIWS	SWISSPROT	ENT CLINANI	TIN TIN	1 HOL	EST LINAN	ZWOL TO	ES L TOWAN	H		SWICEPPOT	ST HIMAN	ST HIMAN	NT		SWISSPRO!			EST HUMAN	Z		ES! HUMAN		
ıß	Most Similar (Top) Hit Top Hit Acession BLAST E No.	4 0F-03 AW 199426 4	4 0E-03 013808	4 0F-03 AFORDRER 1	4.0E-03 A.1041742 4	1.00 00 10.1	4.0E-03 AW103719.1	4.0E-03 AA772898.1	4.0E-03 AF005859,1	03 AF169825.1			3 At 133871 1	3 U22180 1	03 AW500572 4	03 BE548453 1	3 AAB13222 4	T	4.0E-03 AI 183278 2	T	T	3 AI681483.1			4 0F-03 COTTO2	,	32087	A IEGODO A	A 1 482200 0	ALTONZUB.2	ML1032/8.2	1130004. I	T	
	Expression (T	0.97	0.73	0.73	1.95				1.8	23.91	2.48	1.74		L	L			L	L	L	١.		L	0.74 4	4			80 8				L		
	ORF SEQ ID NO:	28932	28008	29300			30383		30625	30774	31421	31423	31510	-	31888	31967	32347	32632	32470	32471	32851	33091	33093		33686	33767	33928	34448		34635	35679	38135		
	Exon SEQ ID NO:	16277	16365	16659	16725		17776	1/832	18002	18118	18499	18502	18583	18774	18918	18991	19336	19594	19452	19452	19787	20013	20015	20101	20538	20842	20797	21308	21479	21489	22478	22923	23112	
	Probe SEQ ID NO:	3521	3612	3909	3977		5057	47.10	5	5314	2705	5708	5792	5993	6140	6217	6672	6677	6970	6970	7098	7331	7333	7424	7843	7947	8103	8614	8787	8797	Ŀ	L	l I	:

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					5	בישום בישוון ומדם פולוום	
Probe SEQ ID NO:	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
11784	24374		1.48	3.0E-03	3.0E-03 AW294812.1	П	UI-H-BI2-ehi-d-06-0-UI.s1 NCI_CGAP_Sub4 Homo sapiens cDNA clone IMAGE:2728842 3
11927	25196		2.86	3.0E-03	3.0E-03 AI525056.1	EST_HUMAN	promrna-5.E07.r bytumor Homo saplens cDNA 5
1,00	<u>L</u>	37056		3.0E-03	1	EST HUMAN	ot77b10.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1022/79 3 similar to contains L1.t3 MER26 repetitive element;
12016				3.0E-03			Homo sapiens gene for CMP-N-ecetylneuraminic acid hydroxylase, partial cds
12190	1	31066		3.0E-03		LN.	Raftus norvegicus mRNA for connexin36 (cx36 gene)
502	L				Q04652	SWISSPROT	RING CANAL PROTEIN (KELCH PROTEIN)
502	ı				2.0E-03 Q04652	SWISSPROT	RING CANAL PROTEIN (KELCH PROTEIN)
768	1				2.0E-03 T70874.1	EST_HUMAN	yd15h03,r1 Soares fetal liver spleen 1NrLS Homo sapiens cuina done liviado: 105341 o
1342	L		2.07	2.0E-03	2.0E-03 M20783.1	۲	Human alpha-2-plasmin inhibitor gene, exons 6 and /
1345	14093	26768	1.4	2.0E-03	2.0E-03 AA661605.1	EST_HUMAN	nu86f01.s1 NC _CGAP_AIV1 Homo saplens CLONA clone IMAGE: 1217.030
1354	ı		16.16	l	AF284446.1	LN	Homo sapiens fumor-related protein DRCz (DRCz) gene, complete cus
1473	1] _			2.0E-03 P48509	SWISSPROT	PLATELET-ENDOTHELIAL TETRASPAN ANTIGEN 3 (PE I A-3) (GPZ?) (MEMBRANE GLTCOPRO I EIN SFA-1) (CD151 ANTIGEN)
	1	_				Ė	Homo sapiens procolagen-lysine, 2-oxoglutarate 5-dioxygenase (lysine hydroxylase, Ehlers-Danlos syndrome hydroxylase, Ehlers-Danlos syndrome hydroxylase, Ehlers-Danlos syndrome hydroxylase, Ehlers-Danlos syndrome
1506	14252	26938	1.84	2.0E-03	455/830 INI	Z	(you) (1000) III was a second of the horizontal and the surface Enjets Danjos syndrome
1506	14252	26939	1.84	2.0E-	4557836 NT	TN	Homo sapiens procolagen-tyeine, z-oxoglutarate protoxygatase (tyeate riyar oxytase, prince prince oxytase) (PLOD) mRNA
1586	1_		6.31	20E-03	03 P29400	SWISSPROT	COLLAGEN ALPHA 5(IV) CHAIN PRECURSOR
1764	L	27207	1.13	_	03 AA450138.1	EST HUMAN	2x42a10,r1 Soares_total_fetus_Nb2HF8_9w Homo Sapiens CDINA clone ilwiAuE.709114.5
1872	14610		1.01	L	2.0E-03 BE144908.1	EST_HUMAN	CM2-HT0183-061099-018-d03 H10183 Homo sapiens cUNA
1988	L	27445			AF302691.1	NT	Mus musculus myelin expression ractor-3-like protein gene, partial cus
2247	1		1.16	L	03 AL163302.2	NT	Homo sapiens chromosome 21 segment HS21C102
2558	l		4.01		2.0E-03 AF187974.1	L	8 Homo sapiens concentrative nucleoside transporte (CN I 1) gene, excri 12
2558	15272				AF187974.1	N	8 Homo sapiens concentrative nucleoside transporter (Livi 11) gene, expr. 12
2584	L	١.		L	03 AW137782.1	EST_HUMAN	UI-H-BI1-adi-g-10-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens CUNA cione III/AGE: 27 1/010 3
24.	1	28818		L	AA450138.1	EST HUMAN	zx42a10.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA cione IMAGE: / 09114 3
25.	1				BF568955.1	EST_HUMAN	602183960T1 NIH_MGC_42 Homo saplens cDNA clone IMAGE-43000 /0 s
	L						H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14
3657	16410	29048	6.62	. 2.0E-	03 X87344.1	ΤN	genes
4093	L	29461		2.0E-	03 P03374	SWISSPROT	ENV POLYPROTEIN (CONTAINS: COAT PROTEIN GF39)
4195	16936		11.03	3.0E	03 U68491.1	LN LN	Rattus norvegicus 5-hydroxydyptamine/ receptor gene, par utal cus
4393	17130		1.12	2.0	:-03 AW297380.1	EST HUMAN	UI-H-BW0-gir-g-03-0-UI-S1 NCI CGAP Judo Homo saprans duna cione ilwa-ce-iz 304 to 3
4397	17134	29765	5 0.97	2.0E	A1064746.1	EST_HUMAN	HA0507 Human fetal live CUNA library monto septens cuna
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Page 180 of 536 Table 4 Single Exon Probes Expressed in Brain

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Droscohija melanogastar shortsintitod closes 2 /ohn)	Drosophila melangaster shortsinhted cless 2 (shs) mRNA, complete cos	yo45e02.s1 Soares adult brain N264H855Y Home saniens c-DNA clone IMAGE 180son 2	Homo sapiens X-linked anhidratic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat recions	801883004F1 NIH MGC 7 Home sealess child stone MA CE 2007750 F1	601876385F1 NIH MGC 55 Hama sapians child clima IMACE: 140.4602 21	Homo sepiens mRNA for KIAAAASa orolein madiel ode	Xanopus laevis xefillin mRNA, complete cals	ATP-DEPENDENT NUCL FASE SITE INIT R	ATP-DEPENDENT NUCLEASE SUBLINIT B	CARBONIC ANHYDRASE-RELATED PROTEIN 2 PRECI IRSOB / CABB 31/CA BB IN /CA SI	CARBONIC ANHYDRASE-REI ATED PROTEIN 3 PRECIDENCE (CARD ALCAND IN CALD)	601887434F1 NIH MGC 17 Homo sapiens CDNA clone IMAGE-41214AB R	ADAM-TS 7 PRECURSOR (A DISINTEGRIN AND METALLOPROTEINASE WITH THROMBOSPONDIN MOTHES 7) ADAMTS-7) ADAMTS-7) ADAMTS-7	AV709075 ADC Homo sersions cDNA class a DOAFFoo El	Lesculentum mRNA for hard-tRNA synthetese (1 v=DS)	wu36h09.x1 Soares_Dieckgrade_colon_NHCD Homo sapiens cDNA clone IMAGE:2522177 3' similar to	2/3311 s1 Spans (ata) liver sales ANETS S4 U.	Geendhabdiús elegans mRNA for galestin I EC 41 complete sels	Homo sapiens lipoma HMGIC fusion partner // HFP) mBN A	Homo sapiens lipoma HMGIC fusion partner (I HFP) mRNA	CM4-BT0366-061298-054-d01 BT0366 Homo saplens cDNA	qm89d11.x1 NCI CGAP 1.u5 Homo saniens 40 NA close 1400 case 21	V477g10.r1 Soares fetal liver spleen 1NEI S Home saniens cONA close MAACE-144200 E	PROTEOGLYCAN LINK PROTEIN PRECURSOR (CARTILAGE LINK PROTEINVI P)	hf37b08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2934035 3' similar to TR:Q60976 Q60976 JERKY.	yx42g06.s1 Soares melancoyte 2NbHM Homo sapiens cDNA clone IMAGE:284442 3' similar to contains L1.b2 L1 repetitive element ;
gle Exon Prof	Top Hit Database Source	L	LZ.	EST HUMAN	L	EST HUMAN	EST HUMAN	N	N.	SWISSPROT	SWISSPROT	SWISSPROT	Т	L	SWISSPROT	T	T	NAM	Τ	Т			т-	Т	EST HUMAN	\mathbf{T}	EST_HUMAN O	EST_HUMAN_L
	Top Hit Acession No.	03 L42512.1	03 L42512.1	03 R87773.1	03 AF003528.1	03 BE798380.1	2.0E-03 BF241410.1	Г	Π	Γ	l			2.0E-03 BF308187.1		T	2.0E-03 X94451.1		2.0E-03/AA677831.1		5031864 NT	5031864 NT	2.0E-03 BE067986.1		ĺ		AW592004.1	N20287.1 E
	Most Similar (Top) Hit BLAST E Value	2.0E	2.0	2.0E-	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03 /	2.0E-03	2.0E-03	2.0E-03	2.0E-03 N
	Expression Signal	2.2	2.2	1.84	2.57	-	1.38	2.08	1.86	3.58	3.58	2.17	2.17	7.5	244	96.0	1.62	1.16	0.61	1.62	9.0	9.0	3.65	0.58	0.8	1.18	1.96	6.07
	ORF SEQ ID NO:	29877			30290	30427	30893	31245	31333	31760	31761	32001	32002	32004	32046	32047	32078		32311	30576	32720	32721	32631	32795	32953	33281	33769	33944
	Exon SEQ ID NO:	1		17397	_	17811		í	- (18800	- 1	_		19029	19064	19065	19090	19271	19305	17939	19674	19674	19506	19735	19879	20188	20645	20810
	Probe SEQ ID NO:	4509	4509	4683	4956	5092	63389	5540	5623	6019	6019	6253	82 83 83	6255	6291	6292	6320	6506	6541	6862	6839	6939	984	2 4 4	7183	7617	7950	8118

Page 181 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	yx42g06.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:264442 3' similar to contains L1.b2 L1 repetitive element;	HYPOTHETICAL 32.8 KD PROTEIN C6G9.05 IN CHROMOSOME I	LAMININ ALPHA-1 CHAIN PRECURSOR (LAMININ A CHAIN)	Homo sapiens Retine-derived POU-domain factor-1 (KPF-1), mKNA	Homo sapiens Retina-derived POU-domain factor-1 (RPF-1), mRNA	AU136679 PLACE1 Homo sapiens cDNA clone PLACE1004839 5	Homo saplens ASQL3 gene, CEGP1 gene, C11orf14 gene, C11orf15 gene, C11orf15 gene and C11orf17	gene	Globalis curiculas minuta in caracteristica como por presentación de la como como como como como como como com	MRZ-UM0026-300300-102-102 UM0025 Incide deptents Control	MR2-UM0025-300300-102-102 UM0025 Homo sapiens culvA	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme EZU s (UBE2D3) genes, complete cds	yp86a09.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:194296 3'	washang s.1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:194296 3	Process and the company (CYTOTACTIN) (NEI IRONECTIN) (GMEM) (JI)	TENASCIN PRECURSOR (IN) (HEXABOXCHION) (CTTOTACIN) (NECOLICIAN) (CHENICAL PROPURSOR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GEOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-700) (GP	AZS) (TERMSCHING) (TING)	BE IA-GALACTOURDAND TARCONAGE (INC.)	BE I A-GALACTO SIDAGE TRECONSON (LANCE)	mo eaplens caspase recruimen donain-containing tracer (2011) years complete eds	Homo sapiens caspase recruitment domain-comaining protein (DOLIN) gene, compress con	QV3-CI 0064-060400-144-601 O L0094 name salvens color	2310a06.51 NCI CGAP GCB1 Hamo sapiens culture livrace. Control of the control of	RC1-C10251-141099-012-d01 C10251 name suprems contra	RC1-CT0251-141099-012-d01 CT0251 Homo saplens cDNA	Human dystrophin gene	PROTEOGLYCAN LINK PROTEIN PRECURSOR (CARTILLAGE LINK PROTEIN) (L.)	RC3-BT0333-310800-116-g04 BT0333 Homo sapiens CDNA	H. sapiens variable number tandem repeat (VN I K) locus UNA	tyesho3 x1 NCI_CGAP_Kld11 Homo sapiens cDNA clone In/AUE: 2265899 3 Similar to 5 W. YA I C_IMPAGE. Q26532 VACUOLAR ATP SYNTHASE SUBUNIT G ;	Homo sapiens SEL1L (SEL1L) gene, partial cds	
and I love	Top Hit Database Source	EST_HUMAN L1.	SWISSPROT HY	SWISSPROT LA			T_HUMAN		NT		٦	EST HUMAN ME	H .	T HUMAN	Γ	T		Т	٦	ISSPROT		Т	٦	П		L HUMAN	П	٦	HUMAN	Į.	EST_HUMAN Q	Г	7
25	Top Hit Acession No.	-			F005855 NT		2.0E-03 AU136679.1			2.0E-03 AB035256.1		AW796111.1	A E224680 4	03 H50832.1	1150000 4	23 H50832.1		03 P24821	03 P48982	03 P48982	03 AF097732.1	03 AF097732.1	AW884269.1	AA251376.1	AW361176.1	2.0E-03 AW361176.1	03 M86524.1	-03 P07354	03 BP330909.1	-03 Z11740.1	03 A 1625745.1	A E457540 2	Z.UE-03/AF 13/310.2
	Most Similar (Top) Hit BLAST E Value	2.0E-03	2.0E-03 Q92350	2.0E-03 P19137	2.0E-03	2.0E-03	2.0E-03		2.0E-03	2.0E-03	2.0E-03	2.0E-03	000	2.0E-03	Lo	2.0E-03		2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	2.0E-03	20.0	∠.∨⊏~∨
}	Expression Signal	6.07	0.54	1.19	0.81	0.81	0.86		0.91	0.54	0.74	0.74	700	40.0		88.0		3.46	1.38	1.38	0.53	0.63	0.81	5.75	0.45	0.45	2.97	2.58	2.14	9.1	3.23		2.41
	ORF SEQ ID NO:	33945	33987	34015	34069	34070	34097			34400					l	35277		35178		35405		Ì	35668		36377		١	33281		37446		١	37807
	Exon SEQ ID NO:	20810	20856	20878	20033	2002	20958		21011	21262	18419	18419		- 1	-1	22104		22008	22219	22219	١	ı	ì	ı	L	1.	L	L	L	L		1	24487
	Probe SEQ ID NO:	4,1	8163	8184	0220	02730	28,28		8318	8570	9094	9000	3	9139	3450	9426		9468	9298	9996	9623	9623	9815	9942	10508	10508	10938	11470	11531	11538	4	200	11926

Page 182 of 536 Table 4

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| Top Hit Descriptor | oy43g08.s1 Soares, parathyrold, tumor, NbHPA Homo sapiens cDNA clone IMAGE:1668634 3' similar to | Complies described to the Control of | AVROZOGA CKY U | H. seplens M1 gene for miscephic each/choling constant | ta6602.x1 Scarss_total_fetus_Nb2HF8_9w Homo sepiens cDNA clone IMAGE:2049051 3' similar to contains Alu repetitive element: | Homo sapiens MSH55 gene, partial cds; and CLIC1, DDAH, G8b, G8e, G6b, G8d, G8e, G8f, BAT5, G5h | JOSKZB, BA 14, G4, Apo M, BAT3, BAT2, AIF-1, 1C7, LST-1, LTB, TNF, and LTA genes, complete cds | AV69/966 GKC Homo sapiens cDNA clone GKCGXD05 5' | Pased8.11 Scares, pineal, gland, N3HPG Homo sapiens cDNA cione IMAGE;232334 6;
as70b08.17 Baretsead colon HPLRB.7 Homo sapiens cDNA cione IMAGE;2334039 3; similar to TR:Q1382;
O44825 411 Binkning con commissions
 | STOORS AND Easted colon HPLRB7 Home septens CDNA clone IMAGE:2334039 3' similar to TR:Q1382: | 413623 AU-BINDING PROTEIN/ENOYL-COA HYDRATASE.; | wkseaus,x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE;2422258 3/

 | WASSELLY, NO. COMP. Mel'16 Hamo sapiens cDNA clone IMAGE:2661242 3' | HORD Sarlang SCI Accolusts Weight FORM OF MYOSIN ((HMWM)) | Tomo saplens mRNA for KIA 41991 models models | CARBONIC ANHYDRASE VI PRECURSOR (CRBBONATE DEHYDRATASE VI) (CA-VI) (SECRETED - 2ARBONIC ANHYDRASE) (SAI IVARY CRBONAIC ANHYDRASE (SAI IVARY CRBONAIC ANHYDRASE)

 | SARBONIC ANHYDRASE VI PRECURSOR (CARBONATE DEHYDRATASE VI) (CA-VI) (SECRETED | AND SOCIETY SALIVARY CARBONIC ANHYDRASE) | Total September 5 VM I gene for synaptic vesicle monoamine transporter, exons 14, 15 | CBAP1D4909 Pediatric pre-B cell acute lymphoblastic leukamia Baylor-HGSC project≔TCBA Homo | Appens conne I CBAP4909
 | Addition nationals elegans spliced leader RNA (SL3 alpha), (SL4), and (SL6) genes | MACOLAN COMES, USUS INT I Homo sapiens cDNA clone IMAGE: 1640262 3' | MO-HT0339-200400 040 Bog LT2002 1
 | APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV) |
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 | 1.0E-03 A | 1.0E-03 AI | 1.0E-03 BI
 | 1.0E-03 O46409 |
| Expression
Signal | 2.41 | 4.37 | 2.99 | 1.76 | 1.48 | 5 | 2,65 | 1 28 | 231
 | 2.31 | 3.78 | 1.78

 | 3.38 | 12.13 | 1.37 | 2.21

 | 221 | 1.65 | 1.28 | 4.05 | 28.0
 | 1.68 | 1.68 | 4.29
 | 7.24 |
| ORF SEQ
ID NO: | 37810 | | | 31050 | | | | 25860 | 26248
 | 26249 | 26489 | 26510

 | 27486 | 27614 | 28383 | 28699

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 | 30127 | 30128 |
 | 30348 |
| | | | | ┙ | 26224 | 24882 | 26178 | 13215 | 13581
 | 13581 | 13831 | 13851

 | 14756 | 14880 | 15745 | 15949

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 | 17505 | 17505 | 17506
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| Probe
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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein Kinase I (CAMKI), creatine transporter (CRTR).	Himpo Jose II	Human class III accord denydrogenase (ADH5) chi subunit mRNA, complete cds	Thermotoce pennelling of the Complete ods	Thermotoga neaponitana alpha-1,6-galactosidasa (aglA) gane, complete cds	BONE PROTEOGLYCAN II PRECURSOR (PG-S2) (DECORIN) (PG40) (DERMATAN SULFATE	NOTE GET CANAIL (DSPG)	Homo saniens transdired hom it is 2 / 15 is 2 / 15 is 2 / 15 is 2 / 15 is 3	ov75f08.x1 Soares_testie_NHT Homo seplens cDNA clone IMAGE:1843175 3' similar to contains MER39.b1 MER39 MER39 repetitive element	ag93f12.s1 Stratagene hNT neuron (#837233) Homo saplens oDNA clone IMAGE:1142083 3' similar to	contains Alu repetitive element;	RC1-CT0279-181099-011-a09 CT0279 Homo saplens cDNA	RCT-C102/9-181099-011-a09 CT0279 Homo sapiens cDNA	QV3-HT0543-220300-130-e03 HT0543 Homo saplens cDNA	W 3612X1 NCI_CGAP_HSC3 Homo sapiens cDNA clone IMAGE:22484463' similar to TR:028195 Q28195 PV41 GFNF	xm72d12.x1 NCI CGAP Kid11 Home confee Chit	AV759949 MDS Home series convenience of the property of	801433087F1 NIH MGC 72 Homo seriors albity also have processed at	AV731520 HTF Hamo septens cDNA clone HTFA 1205 E	to 54/1.x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:2063013 3' similar to contains Alu	HARBOTRE MILL MOC 27 11	ROLCTONE 240000 02 JOS OFFICE TO SEPTEMBERS CON CONTRACT 3872035 5	LINE-1 REVERSE TRANSCROOTASE LICENAL CONTRACTOR	APOLIDODBOTENIA IV BBEZINGS (1805)	Homo sanlens (AVI OT1 2000)	MITOCHONDRIAL RIBOSOMAL DESCRIPTIVARIA	Glycyrthiza glabra GabAS1 mBNA for hata-amin'n circhicas	X. laevis mRNA for C4SR protein
gie Exon Pro	Top Hit Database Source	<u> </u>	LN	Ę	L	LN	TOGGSSIMIS	TN	LZ	EST_HUMAN	$\overline{}$	ESI HUMAN	7	7	EST HOMAN	EST HUMAN	Т	┰	EST HUMAN	_		T	T	SWISSPROT	Т		SSPROT		
	Top Hit Acession No.	03 U52111.2	Ī		-	T	03 001129	1		03 AI024350.1	3 4 4 706303 4	١	1.0E-03 AW36Z393.1	T	T			1				T				45.1		AB037203.1 NT	X96469.1 NT
	Most Similar (Top) Hit BLAST E Value	1.0E-03	1.0E-03	1.0E-03	1.0E-03/	1.0E-03 /	1.0E-03	1.0E-03 /	1.0E-03	1.0E-03	1 00 00	4 60 50	1.0E-03 A	4 20 00 00	0 -0-10	1.0E-03 AI583847,1	1.0E-03 A	1.0E-03 AV759949.1	1.0E-03 BE894488.1	1.0E-03 A	1.0E-03 Al347355.1	1.0E-03 BE780572.1	1.0E-03 AW847341.1	9.0E-04 PC	9.0E-04 Po	9.0E-04 AJ		9.0E-04 AB	8.0E-04 X9
	Expression Signal	0.58	3.68	3.68	1.66	1.66	0.81	0.65	0.75	1.72	0 46	70	1 70	97.6	21.3	3.29	1.36	3.05	4.48	1.38	1.98	7.05	1.37	0.7	1.28	9.0	98.0	1.42	1.02
	ORF SEQ ID NO:		34858			35341	35570	35929		36076		36495	36498	38580			36727		37805			30612	30809	30465			32185	+	
	Exon SEQ ID NO:	_	ii		22160		22371	22711	22718	22862	23149	23259	23269	23342		23413	23491	23776	24470	24616	25342	25365	25187	17848	18391	18942	19164	22198	14218
	SEQ ID NO:	8978	9017	9017	9507	9507	9720	10063	10068	10214	10503	10563	10563	10651		10726	10808	1108	11906	12124	12371	12478	12821	633	6596	6165	6395	9543	14/1

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Table 4
Single Exon Probes Expressed

			T	;	T	T	Γ	Z	Т	_	Т	П		Т		Т		Г	г	ГТ.	_		Τ,	_				_
Single Exon Probes Expressed in Brain	Top Hit Descriptor	W78g11.X1 NCI CGAP 1 110 Home	repetitive element	HYPOTHETICAL 30 3 UP BEC. (2.5)	QV0-CT0225-02009-030-407 CT0205-11	nk27e11.s1 NCI_CGAP_Co11 Home saplens cDNA Access 144.05	repetitive element ADAM-TS 7 PRECURSOR (a DISINITE CENTRE CENTRE CENTRE CENTRE ALI	MOTIFS 7) (ADAMTS-7) (ADAM-TS7)	Bos taurus micromolar calcium activadad noutral	2033b08.r1 Stratagene colon (#837704) Home series 1 (CAPN1) gene, exons 11-20, and partial cds	Gorilla gorilla Involucrin gene medium aligie, complete color	dd13f08.x1 Soares_placenta_8tx9weeks_2NbHP8tx9W Homo sepiens cDNA clone IMAGE:1723819 3' similar to gb:X61602_cds1 VASCULAR ENDOTHELIAL GROWTH FACTOR RECEDED.	obsection CCAP GCB1 Local Control CCAP GCB1 Local CCAP GCB1 CCAP GCB1 Local CCAP GCB1 CCAP GCB1 Local CCAP GCB1 CCAP GCB1 CCAP GCB1 CCAP GCB1	MER22 repetitive element;	ajS6h03.s1 Soeres_testis_NHT Homo sapiens cDNA clone IMAGE:1394357 3' KK2745F Human fetal haari amhda 7AD E.	REPETITIVE ELEMENT	BIFUNCTIONAL ENDO-1,4-BETA-XYI ANASE XYI X BEFAITESER	XS08e02.X1 NCI_CGAP_Kid11 Homo sapiens cDNA class 144 SE 2323	Human familial Alzheimer's disease (STM2) gene complete de	DKFZp588M2024_J1 588 (synonym: hute1) Homo saplens cDNA clone DKFZp588M2024	30s taurus micromolar calcium anthinated manifest	mf15h02.s1 NCL CGAP_pr1 Home septems cDNA clare 11.00 gene, expns 11-20, and partial cdo	Haemophilus Influenzae Rd section 63 of 163 of the complete genome	8870b08.x1 Barstead colon HPLRB7 Homo saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens cDNA clone IMAGE:2334039 3' similes to TBI Anna Saplens clone IMAGE:2334039 3'	8870b08x1 Barstead colon HPLRB7 Home Control Part ASE.	Q13825 AU-BINDING PROTEINENOYI-COA HYDBATASE	RC3-CT0254-130100-023-f01 CT0254 Homo sapiens cDNA	Homo saplens chromosome 21 segment US 21 Onza
igle Exon Pr	Top Hit Database Source		EST HUMAN	SWISSPROT	EST_HUMAN		ESTAUMAN	SWISSPROT	Ν̈́	EST_HUMAN	Ā	EST HIMAN	Т	EST_HUMAN	$\neg \vdash$	EST_HUMAN	╗	HOMAN	П	ESI_HOMAN		T_HUMAN	1	EST HUMAN O		7	HOMAN	
ਨੋ	Top Hit Acession No.		6.0E-04 AI817088.1	010341	5.0E-04 AW851844.1	5.0F-04 A45.48031 4		T		-	-04 MZ3004.1	04 A1188382.1		04 AA814519.1 E	-			T	T			T	Z/40.1	AI720263.1 ES		AMZESSES.1 ES	T	
	Most Similar (Top) Hit BLAST E Value		6.0E-04	5.0E-04 O10341	5.0E-04	5.0F-04	L	2.05	3.0E-04	3.0E-04/	2.0E-04	5.0E-04	L	3.0E-04 A	200	5.0E-04/N83788.1	S OF OA	5 OF OA 11	5 0E-04 AT 048507 0		5.0E-04 AF248054.1	4 OE-04 AA568513.1		4.0E-04 AI7		4.0E-04 AIV	4.0E-04 AL	
	Expression Signal	100	1.04	10.0	1.4	1.35	232	000	7 80	3.75		5.2	80	300	89 0	1 44	4.1	0.48	2.38	3	4.08	148		1.79	1 70	3.18	1.81	
	ORF SEQ ID NO:		28051			28815	29096	30877	32303	33037		33677	34033	35013	35195	35268	35344	-		30877	+	28076		26267	26268	26880	27538	
	SEQ ID NO:	25058	1	1	1	16166	16457	18186	1	19960		20551	20896	21847	22025	22098	22162	22825	23577	18186	25184	13435	_	13597	13597	14196	14807	
	Probe SEQ ID NO:	12818	939	1490		3408	3704	5386	6532	7276		7856	8202	9177	9271	9418	9509	10177	10897	11713	1	1 1		827	827	1449	ı ,	
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					3.10	Single Exoll Flores Explosed	
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
200	14860		-	4.0E-04	AL046704.1	П	DKFZp434D059_11 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434D059 5
2833	1	28088	2.21	4.0E-04	096615	ISSPROT	SERICIN-2 (SILK GUM PROTEIN 2)
3162	1			4.0E-04	AF281074.1	Ę	Homo sapiens neuropunt 2 (1414 2) gene, company to the lima GE:951930 3' similar to gb:M21121 T-CELL
900	ļ	20853	3.18	4.0E-04	AA576331.1	EST HUMAN	SPECIFIC RANTES PROTEIN PRECUENCE (HUMAN);
4209				4.0E-04	AA576331.1		nh10a10.s1 NCI_CGAP_Cof Home sapiens dury durie invoc
4500	L			4.0E-04	AA086324.1	T	zn61c08.s1 Stratagene muscle 93/209 fruit septems control of the c
5028	L		3.1	4.0E-04	BE560660.1	EST_HUMAN	601343690F1 NIT MICC. STICKED STRONG RECEPTOR PRECURSOR (CASR) (PARATHYROID CELL
7468	1		13	4.0E-04	P48442	ISSPROT	CALCIUM-SENSING ACCOUNTS.
7434	L	L			4.0E-04 AL161566.2	7	Arabidopsis thattana DNA chromosonie 4, caring ingeneral control of the control of the chromosonie 4, caring chore MAMMA1001620 5'
7618	١	33394			4.0E-04 AU122079.1	٦	AUTZZU 8 MANINA I HAITO SEPANO SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CON CONTROL SENIENS CONTROL SENIENT CONTROL
8434	L				4.0E-04 BF240712.1	Т	6018/0963-1 NIH_MICC_SO FIGURE SEPTIMES CON CONTRACE:284142 5
8442	L	1 34270			4.0E-04 N25507.1	Т	MASSELL, I Scales included to the sapient of the IMAGE:16443413'
9590	L	3 35426			4.0E-04 AI025699.1	HOWAN	Mis misculus neuroplin-2/a17) mRNA, alternatively spliced, complete cds
9740	22391		1.22		4.0E-04 AF022855.1	2	From some SMARCA4 isoform (SMARCA4) gene, complete cds, alternatively spliced
12380	L		2.05		4.0E-04 AF254822.1		nd no sable is Own (Section 1) Amount ham 2) Homo saplens cDNA clone DKFZp761.1221 5
15.	L	7 25608			3.0E-04 AL119426.1	EST HUMAN	ARICE TO SECRET ORY PHOSPHOLIPASE AZ RECEPTOR PRECURSOR (PLAZ-R)
190	13003				3.0E-04 P49259	DRACCIMO	Liman short chain and CoA dehydrogenese gene, exons 1 and 2
88	0 13629				3.0E-04 U83991.1	IN TOL	AZ28AR3 V NCI CGAP Kid11 Homo saplens cDNA clone IMAGE:2028197 5
183	1 14570	0 27282			AI262100.1	TO TOWN	#23an2 v1 NCI CGAP Pr28 Homo sapiens cDNA clone IMAGE:2119082 3
1846	6 14584				3.0E-04 Ai399674.1	TOUR HOMAN	INTERNALIN B PRECURSOR
3303	3 16064	4 28712			3.0E-04 P2514/	DWING TOT	1258014 rt Soares fetal liver spleen 1NFLS_S1 Homo sapiens cDNA clone IMAGE:446478 5
3308	8 16068				3.0E-04 AAZU3342.1	TOGGGGWG	IGI LITAMATE DEHYDROGENASE 2 PRECURSOR (GDH)
3946	18696	6 29335		3.0E-(3.0E-04 P49448	DI PONTO	Home sablens Xg pseudoautosomal region; segment 1/2
<u>දූ</u>	16779	6.	1.33	3.0E-(34 AJ2/1/35.1	IN TOU	PCO-HT0014-310599-028 HT0014 Homo sapiens cDNA
4072	72 16816	8	1.12	3.0E-(34 BE140609.1	EST HOMAN	NOCH H0214 010000 007-012 HT0339 Homo saplens cDNA
4766	<u>L</u>	8	4.72	3.0E-	04 BE153778.1	EST HOMAN	PM04T1 0555-150250 058-009 DT0045 Homo saplens cDNA
482	17558	30180	0.95	3.0E-	04 AW837723.1	HOLINAN	AND STANCE CRAP Lut Home sapiens cDNA clone IMAGE:1143328 3'
5063	33 17782	30389		3.0E-	04 AA613145.1	HOWEN TOWAR	Home satisfies chromosome 21 segment HS21C081
6052	52 18832	32	7.86	3.0E-	04 AL 163281.2	N I	Homo soniens chromosome 21 segment HS21C078
6722				3.0E	3.0E-04 AL 163278.2	N I	PROTEIN: TYROSINE PHOSPHATASE DELTA PRECURSOR (R-PTP-DELTA)
7491	31 20163	33258	56 0.84	╝	4 P23468	SWISSERS	

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	FIBRORI AST OROWITH EACTORS OF SECTION OF THE SECTI	248408.r1 Soares, lestis, NHT Homo sepiens cDNA clone IMAGE:795471 6' similar to ab:M62762	WACCOLLAR A I P SYNTHASE 16 KD PROTEOLIPID SUBUNIT (HUMAN);	424905.s1 Soares_tests_NHT Homo explens cDNA clone 1391288 3' similar to gb:M36072 60S	CITAMIC ACID BIOLET BOOK (HUMAN);	Hone sanless administration of the s	nc38e04.r1 NCI_CGAP_Pr2 Homo septens cDNA clone IMAGE:1010430 similar to containe L1.12 L1	Induction element	riumo sapiens mikina for KIAA0749 protein, partial cds	UNT-25947-138 _T 547 (synonym: hfbr1) Homo sepiens oDNA clone DKFZp647L185 6. Homo sepiens SCG10 like-protein, helicase-like protein NHL, M88, and ADP-ribosulation factor related	protein 1 (ARFRP1) genes, complete cds	AU146/0/ HEMBB1 Hamo capiens oDNA clone HEMBB1001263 3	Human dystrophin gene	Human dystrophin gene	4h98e11.x1 Soares_NFL_T_GBC_S1 Homo saplens oDNA clone IMAGE:1855052.3' similar to contains	Horoconia Copenio Benefit	Mile miscrific 6' flooder 21 segment HS21C003	machines of learning region of Prox gene	Human germline T-cell receptor bela chain TCRBV475444T TCBCA-00-1	TCRBV16S1P, TCRBV16S1, TCRBV11S1A1T, HVB relic, TCRBV28S1P, TCRBV34S1, TCRBV34S1	TOMBYSS1, ICRBV4S1A1T, TRY4, TRY6, TRY7, TRY8, TCRBD1, TCRBJ1S1, TCRBJ1S2.>	Britoscus XI Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539760 3'	From Sapiens would, beta, 4 (TUBB4) mRNA	4 7 2 B 1 0030 - 1 000 B 1 0036 Hamo sapiens cDNA	Dhesedite in the second of the	Mid-att of Section 11 to 1 to 1 to 1 to 1 to 1 to 1 to 1	May 1911, 11 Scales, plined, gland, N3HPG Homo sapiens cDNA clone IMAGE:232556 6	Gallus gallus protessome 29 PD 2011 11 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	ganto processorire zo Nut subunit ndmolog mKNA, complete cde Danio rerio hagoromo gene, exons 1 to 6, partial cds
igle Exon Pri	Top Hit Database Source	SWISSPROT	For	EST HUMAN	EST HIBAN	SWISSPECT	NT	FOT LINAM	NAME OF LAN	EOT LILITARI	NAMON I	N - 100	TO HOMAN	Z!	Z	EST HUMAN	LN	Z			Ŀ	ECT LINVANI	NAMOR 10-	EST HIMAN	EST HIMAN	LN FN	T HI IMAN	Т	Т	
Sir	Top Hit Acession No.	P22607	3.0E-04. AA454055.1	3.0E-04 AI892139.1	-04 AA781201 1	P13816	4501960INT	AA228301 1	3.0E-04 AB018292 1	3.0F-04 A 134483 1		T	T	2.0E-04 M86924.1	Τ	-04 AI286021.1	04 AL 163203 2	Γ			04 168081 1	-	74738	3	2.0E-04 AW978441 1	Τ	Τ		Γ	
	Most Similar (Top) Hit BLAST E Value	3.0E-04 P22807	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3.0E-04	3 OF -04/	2000	2.0E-04/	200	205-04	4.0E-04	2.0E-04	2.0E-04	2.0E-04 A			2.0F-04	2.0E-04 A	2.0E-04	2.0E-04 B	2.0E-04 A	2.0E-04 U	2.0E-04 H	2.0E-04 H	2.0E-04 U09226.1	2.0E-04 AE
	Expression Signal	3.23	1.34	0.65	3.73	0.54	1.38	4.81	3.08	2.75	2.65	8	10 71	10 74		3.93	2.18	1.12	<u> </u>		4.47	1.1	1.1	1.99	0.79	4.93	1.74	1.74	1.63	1.1
	ORF SEQ ID NO:	33983	35673		36220	36367	37466	30617	30818		25624	25892	26324	26325							28033	28398	28740	28834	29282		29990	29991		30324
	SEQ ID NO:	20851		22726	23003	23141	24164	25386	25230	25000	12984	13251	13656	13658		13911	13917	14563	_		15295	16762	16088	16186	16642	16864	17355	17355	17474	17721
	Probe SEQ ID NO:	8157	0850	10078	10356	10495	11555	11976	12338	12730	121	468	887	887		1158	1163	1824		<u>—</u>	2581	2986	3328	3429	3892	4122	4620	4620	4742	4898
																			_					_						

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	Top Hit Descriptor	MYOSIN HEAVY CHAIN, SMOOTH MUSCLE ISOFORM (SMMHC)	AV654352 GLC Homo sapiens cDNA clone GLCDUH10 3'	te03b11.x1 NCI_CGAP_U3 Home saplens cDNA clone IMAGE:2207709 3'	EST11191 Uterus Homo sapiens cDNA 5' end similar to EST containing O family repeat	Lorno sentense cell cycle procression 3 protein (DN.3) mRNA	Truini Saprais our spin programmer remember of and unknown defe	MILE MUSICULES & FUCUELI CALIFORNIA PEROF OFFICE AND ALMA ADMINISTRATION OF STATEMENT OF STATEME	AU121/12 MAMMAT Home septems CDNA court Mamma septems CDNA	QVU-C USS/-18050U-187-810 O 1909/ HOMO! OR	LINE-I REVENUE INVINCENTIAL ASSOCIATED PROTEIN (165 KD CONNECTIN-	MYOMESIN Z (WATRO TEIN) (1951/D 1111/17/2000/17/17/17/17/17/17/17/17/17/17/17/17/17/	Solanum lycopersicum phytochrome F (PHYF) gene, partial cds	Solanum lycopersicum phytochrome F (PHYF) gene, partial cds	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORC1L3, ORC1L4 genes,	complete cds)	Home sapiens DNA, DLEC1 to ORC1 L4 gene region, section 1/2 (DLEC1, ONC) L3, ONC) L4 genes,	complete cds)	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Human immunoglobulin C(mu) and C(detta) neavy critari garies (constant egrens)	al 22a 12.51 Soares Testis Unit I nomo saprems como como como como como como como co	GASTRULA ZINC FINGER PROTEIN ALCOL 28.1	RC3-H10254-151099-011-005 II 10254 II umo amilans cDNA clone IMAGE 742964 51	MINISTER LINE CONTRACTOR AND ACTOR HTEADAM 5	AV (303/3 TILT TIGHT) September and expression of the State of the Sta	Figure Saprens par use of 1141 Ecopies gards, come in AGE:2140269 3' similar to contains Alu repetitive	goninati	UIH-BI1-adm-c-04-0-UI.st NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2717190 3	vb79b10.x5 Stratagene ovary (#937217) Homo sapiens cDNA clone IMAGE:773713'	RETROVIR IS REI ATED POL POLYPROTEIN ICONTAINS: REVERSE TRANSCRIPTASE;	ENDONUCLEASE]	UI-H-BIG-aab-e-09-0-UI-s1 NCI_CGAP_Sub1 Homo sapiens cunA cione IMAGE:2708825 3	1十七日での中央の中央の一名の一名の一名の一名の一名の一名の一名の一名の一名の一名の一名の一名の一名の
Siligle Exoli Plobes Explosed	Top Hit Database Source	SWISSPROT	Γ	Т	Т	NAMOR		T	П	Т	SWISSPROI	SWISSPROT	NT N	NT	_	NT		N N		П		٦	Т	Т	HOMAN	Z	EST HUMAN	Г	T	T		Т	EST_HUMAN
	Top Hit Acessian No.	P35748	AVERAGES 4	A IRROPARO 4	A A COCCETA	AAZ90002.1	4758179INI	AF140708.1	4 AU121712.1	2.0E-04 AW860963.1	P08548	2.0E-04 P54298	U32444.2	2.0E-04 U32444.2		2.0E-04 AB026898.1		2.0E-04 AB026898.1	2.0E-04 AF020503.1	2.0E-04 X57331.1	AA725700.1	2.0E-04 P18715	2.0E-04 BE149303.1	AA405777.1	20E-04 AV730373.1	AJ243213.1	2 DE-04 A1440282 1	1 AM 138740 1	Z.OE-04 ANY 1307 43.1	+ MIOZ 1304. I	1.0E-04 P11369	34 AW013847.1	04 AW013847.1
	Most Similar (Top) Hit BLAST E Value	2.0E-04			Z.UE-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04		1	1	ı	2.0E-04		2.0E-04						- 1				١		1		1.0E-	1.0E-
	Expression Signal	104	5	0./3	C/.L	96.0	0.81	0.59	2.6	0.55	14.88	1.42	1.06	1.06		1.23		1.23	1.96	0.49	0.49	0.6	1.19	1.77	5.23	1.61			2.80		3.3	,	4.74
	ORF SEQ ID NO:	30424	2770	31146	31160	31373	31584	31892				33295				34012	l	34013			35086			35761	36684			1	3/350		26472		26509
	Exan SEQ (D NO:	47000	8	18256	- 1			1	19818	19910	20191	l	20550	1	1	20876	ı	20876	l	L	21912	1	1	22565	23440	3 23796	i	_1	⅃	24441	13812	ı	1
	Probe SEQ (D NO:	0000	8	5457	27	5664	5857	6144	7130	7225	7520	7530	7855	7855	3	8182		8182	8463	8642	9233	8319	9876	9946	10755	11128		112/0	11403	11857	1053	1092	1092

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		7	\top	\neg			100	_			\neg	_	_	_	-		٠,	_	_		_,	_							
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Anguilla anguilla dopamine D1A1 receptor (d1A1) مصه حمساطه حماد	Kaposi's sarcoma-associated herpesvirus ORF 68 gene, partial cds; and ORF 69, kaposin, v-FLIP, v-cyclin, latent nuclear antigen, ORF K14, v-GPCR, putative phosphoribosyfrormylglycinamidine synthase, and LAMP (LAMP) genes, connicts cds.	Kaposi's sarcome-associated herpesvirus ORF 68 gene, partial cds; and ORF 69, kaposin, v-FLIP, v-cyclin, latent nuclear antigen, ORF K14, v-GPCR, putative phosphoribosyfformylglycinamidine synthase, and LAMP	Lingual garles, compiete cds	SPLICEOSOME ASSOCIATED BEATERING A CAR ASSOCIATED	101f11.x1 NCI_CGAP_Ges4 Homo sapiens cDNA clone IMAGE:2140269 3' similar to contains Alu repetitive	Mouse ginhs 1 thosally collection and buy	AV647727 (I C Home seniors of National Control of	Homo sapiens KIAAA337 2000 51-1-1-1 IVIA Appar	Homo satisfies KIAA0237 cano madina (VIAA0237), mKNA	LINE-1 REVERSE TRANSCRIPTAGE USING SO	753F Head Homo saniens child Actor 723	763F Heart Homo septions of NA clause 7.53	nc02e12.st NCI CGAP Pr3 Home seniors a DNA along MA CE and	nj25a04.s1 NCI_CGAP_A41 Homo septems CDNA close IMAGE:983488 3' similar to gb:M97252 KAI MANN SYNDRAME BECTERN BETCHING COME	av57410 x1 NCI CGAP CAS Home selection (HOWAN); contains Alu repetitive element;	gv67d10x1 NCI CGAP Ov22 Hours septem CDNA The INACE 185683 3	ab94n08 s1 Stratage him /#037340/ Long - Inches in the Control of	W28e08.x1 Soares NFL T GBC S1 Homo septems cDNA clone IMAGE:254464 3:	OYSTATIN-RELATED EPIDIDYAAI SPERMATOCEAUC BROOKING 185	yd72c08.r1 Sogres (stal liver spiego 1/NE) S Homo som long control (character 180)	Homo sepiens phospholinid screen has a 17P SCE31	LINE-1 REVERSE TRANSCRIPTASE HOMOI OG	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	Mouse abha leukocyte interferon gene, complete ade	UI-H-BIT-ann-4-02-0-LII st NCI COAB Substitution	UI-H-BIT-aew-a-02-0-UI:s1 NCI CGAP Sub3 Homo earliens cDNA class (MAOE:2720546 3	Homo sapiens mRNA for KIAA1142 protein, partial cds
igle Exon Pro	Top Hit Database Source	-N	Ę	<u> </u>	12	SWISSPROT	EST HUMAN	L L	EST HUMAN	LN	1Z	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	SWISSPROT	L		SWISSPROT	SWISSPROT	Т	Т	EST_HUMAN	\neg
is	Top Hit Acession No.	5-04 U62918.1	-04 AF148805.1	AF148805.1	1.0E-04 AB048342 4	262203	1.0E-04 AI440282.1	1.0E-04 M14042.1	1.0E-04 AV647727.1	7662015INT	7662015 NT	08547	19615.1	-04 T19615.1	-04 AA177111.1		-04 AI251980.1	Γ				-04 T77153.1	863876	04 P08547		04 M28587.1	24 AW 205336.1		04 AB032968.1
	Most Similar (Top) Hit BLAST E Value	1.0E-04	1.0E-04	1.0E-04	1.00.04	1.0E-04 Q62203	1.0E-04	1.0E-04	1.0E-04/	1.0E-04	1.0E-04	1.0E-04 F	1.0E-04 T19615.1	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04 A	1.0E-04 AI806220.1	1.0E-04 O88969	1.0E-04 T	1.0E-04	1.0E-04 P	1.0E-04 P	1.0E-04 M	1.0E-04 A	1.0E-04 A	1.0E-04 AI
	Expression Signal	3.12	3.25	3.26	2.09	1.06	0.91	2.11	1.15	1.28	1.28	1.49	0.57	0.57	0.95	0.92	15.6	17.82	96.0	2.27	1.46	0.49	1.86	2.74	1	2.13	2.05	2.05	1.76
	ORF SEQ ID NO:		27058	27069	27308	28689	29110	29412	29437	30368	30369	31487	31556	31557	32105	32605	32841	32841	33719	35088	35097		35381		35975		37479	37480	37566
	SEQ ID NO:	14057	14370	14370	14592	16039	16472	16782	16807	17756	17755	18560	18623	18623	19116	19572	19776	19776	50689	21915	21926	21892	12 18 18 18	22727	22763	23971	24166	24166	24246
	Probe SEQ ID NO:	1309	1623	1623	1854	3278	3719	4037	4062	9030	929	69/9	5834	5834	8346	6738	7086	7470	7894	9238	9247	9326	9546	10079	10115	11312	11567	11567	11649
														_								_	_			_		_	_

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	7-	_	_								_									_						_	_		
Top Hit Descriptor	xx49g12.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:28166183'	NEURONAL-GLIAL CELL ADHESION MOLECULE PRECURSOR (NG-CAM)	NEURONAL-GLIAL CELL ADHESION MOLECULE PRECURSOR (NG-CAM)	7729a10.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:3296058 3' similar to contains L1.t3 L1	repetitive element;	ah45c11.s1 Scares_testis_NHT Homo sapiens cDNA done 1292468 3'	QV4-SN0023-070400-166-b04 SN0023 Homo sapiens cDNA	PROLYL 4-HYDROXYLASE ALPHA-2 SUBUNIT PRECURSOR	UI-H-BI1-eer-d-05-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE: 2720289 3'	ULH-BI1-aer-d-05-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2720289 3'	Homo saplens gene for cholecystokinin type-A receptor, complete cds	Homo sapiens methyl-CpG binding protein 1 (MBD1) gene, exon 15b	xx334g05.x1 NCI_CGAP_Br18 Homo sapiens cDNA clone IMAGE:2568728 3° sImilar to contains L1.2 L1 repetitive element:	qv23f06x1 NCI_CGAP_Lym6 Homo sapiens cDNA clone IMAGE:1982435 3' similar to contains element	MIN repetuve element; PROLYT A HYDROXYTASE AT PHA. 2 STIBLINIT BDECTIDEND	THOSE THE DROWN TASE ALT THE SOBONI PRECORSOR	Homo sapiens MSH55 gene, partial cds; and CLIC1, DDAH, G6b, G6c, G6b, G6d, G6e, G6f, BAT5, G5b, GSK2B, BAT4, G4, Apo M, BAT3, BAT2, AIF-1, 1C7, LST-1, LTB, TNF; and LTA genes, complete cds	Pisum sativum mRNA for beta-1,3 glucanase (gns2 gene)	Pisum sativum mRNA for beta-1,3 glucanase (gns2 gene)	Human platelet-derived growth factor A chain (PDGFA) gene, exons only	wy/8a04.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2554638 31	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds	zs88h01.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:704693 3' sImilar to contains Alu repetitive element contains element MSR1 repetitive element	RC3-CT0208-220999-011-E04 CT0208 Homo saplens cDNA	RC3-CT0208-220999-011-E04 CT0208 Homo sapiens cDNA	HUM072014F Human fovea cDNA Homo sapiens cDNA clone EST HFD072014	HUM072014F Human fovea cDNA Homo sapiens cDNA clone EST HFD072014	PROBABLE GLYCEROL3-PHOSPHATE ACYLTRANSFERASE, MITOCHONDRIAL PRECURSOR (GPAT)	Homo sapiens chromosome 21 segment HS21C078
Top Hit Database Source	EST_HUMAN	SWISSPROT	SWISSPROT		EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT		Г	Г	N	EST HUMAN	1	SWICKBROT	7	ĮN.	L	NT	N	T_HUMAN	±N	EST HUMAN	Т	EST_HUMAN	EST_HUMAN	Г	1	П
Top Hit Acession No.	1.0E-04 AW269061.1	203696	Q03696		1.0E-04 BE676399.1	9.0E-05 AA718933.1	9.0E-05 AW866218.1	Q60716	9.0E-05 AW204958.1	9.0E-05 AW 204958.1	9.0E-05 D85606.1	9.0E-05 AF120982.1	9.0E-05 AW073078.1	, of or only	9.0E-05 A1287676.1	2007 10	9.0E-05 AF129756.1	8.0E-05 AJ251646.1	4,3251646.1	8.0E-05 M83575.1	8.0E-05 AW044605.1	M69197.1	8.0E-05 AA279333.1	AW847445.1	7.0E-05 AW847445.1	49075.1	49075.1	122949	7.0E-05 AL163278.2
Most Similar (Top) Hit BLAST E Value	1.0E-04	1.0E-04 Q03696	1.0E-04 Q03696		1.0E-04	9.0E-05	9.0E-05	9.0E-05 Q60716	9.0E-05	9.0E-05	9.0E-05	9.0E-05	9.0E-05	100	9.0E-05 A128/6/	9.00	9.0E-05	8.0E-05	8.0E-05	8.0E-05	8.0E-05	8.0E-05	8.0E-05	7.0E-05/	7.0E-05	7.0E-05 L49075.1	7.0E-05 L49075.1	7.0E-05	7.0E-05/
Expression Signal	2.01	2	2		2.51	2.76	1.14	1.81	9.0	9.0	3.02	2.78	2.68	;	3.5	3	6.63	1.97	2.75	67.0	0.87	1.84	4.65	1.14	1.14	1.1	1.1	4.1	2.99
ORF SEQ ID NO:		37643	37644			26102	27455	31601	33242	33243		35125	37027	200	31601	3		26237			29808	37045		25773	25774	25965	25966	26453	28168
Exen SEQ ID NO:		24319	24319		25203	13457	14733	18660	20149	20149	21951	21953	23752	25.05	18660	3	26259	13574	13614	15718	17184	23769	25242	13138	13138	13337	13337	13793	15431
Probe SEQ ID NO:	11692	11725	11725		12131	682	1997	5873	7478	7478	9376	8278	11082	74.30	11817		12178	802	844	2950	4448	11099	12765	337	337	554	554	1033	2724

Page 192 of 536 Table 4

		T	٦	T	Т	Т	Т	T	Т	_		7				_	7	_	7	-	т-	_	-	_		-	-					
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Dictosfelium discoideum gens for TREA commists ada	Homo sapiens chromosome 21 segment HS24 Coot	Caenorhabdilits elegans Skorto homolog mRNA complete add	Rat cycmegalovirus Masstricht complete genome	Inh93g01.st NCI CGAP Br2 Home septiens child charalitations of	EST04984 Fetal brain. Stratanene frauttonenen Jane Americanian et in in in in in in in in in in in in in	Homo sapiens sarcoglycan, epsilon (SGCE), mRNA	Homo saplens chromosome X open reading frame R (CXOBER) apply	Homo saplens chromosome X open reading frame B (CXORFR) mBNA	wb54h06x1 NCI_CGAP_GC6 Homo sepiens cDNA clone IMAGE:2309531 3' similar to gb:J03260 DNA	H sorions days and dependently),	Histories flow-sorted chromosome 6 Hindlil fragment, SC6pA28B10	Homo sapiens monocytalne through a forting in the contraction of the c	CONTACTIN PRECIPEOD (CI VODBOTE), CASA	CONTACTIN PRECIENCE (ALICOPACIEN GETSS)	W500111 Scares fetel line relicent MHT S. University of the state of t	OROBOTA STATES NEW TORN STORMS TO THE STATES OF THE STATES	RC4-BT0311-141199-011-hos RT0311 Home septions CONA cione invidenci 104588 3	RC4-BT0311-141199-011-h06 BT0311 Homo seniens cDNA	208c08.st Scares_pregnant_uterus_NbHPU Homo sepiens cDNA cione IMAGE:491726 3' similar to	PM4-NN0050-310300-001-410 NN0050 Home seniors - DNA	COMPLEMENT DECAY. ACCEL FRATING FACTOR DEFOLIPSOR	CAB-BINDING PROTEIN PRECURSOR (CARP)	C4B-BINDING PROTEIN PRECURSOR (CARP)	ye28c12.r1 Stratagene Jung (#937210) Homp seniens cDNA clara IMA GE1110062 FT	hi37a03.x1 Soares NFL T GBC S1 Homo sabiens china IMAGE 202444.3	y69d08.s1 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:143535 3' similar to contains Alu	AkSAO2 rd Spares present them shipping element;	MRD-NT0038-250400-004-60 NT5038-U	7028a08 x1 NCI CGAP Bring Home sealers CDNA	QV4-ST0234-241199-040-h11 ST0234 Hamo saplens cDNA
le Exon Pro	Top Hit Database Source	N	N	LN LN	ΡĀ	EST HUMAN	EST HUMAN	1	Z	12	EST HIMAN	LN LN	i i	LN	SWISSPROT	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	HST HIMAN	T	Г	Т	T	Π	Π		Т	Т	Т	П
Sing	Top Hit Acession No.	-05 AB009080.1			9845300	05 AA505582.1	.05 T07095.1	10835046 NT	4885170 NT	4885170 NT	05 AI655241.1	T		-	T		-	-			6.0E-05 AA150482.1			Γ			8.0E-05 AW627985.1 E		Τ	6.0E-05 AW890110.1	T	П
	Most Similar (Top) Hit BLAST E Value	7.0E-05/	7.0E-05	7.0E-05	7.0E-05	7.0E-05/	7.0E-05	7.0E-05	6.0E-05	6.0E-05	6.0E-05 A	8.0E-05/Z	6.0E-05/Z	6.0E-05	6.0E-05	6.0E-05	6.0E-05	6.0E-05/A	6.0E-05 B	6.0E-05/B	6.0E-05.A	6.0E-05 A	6.0E-05	6.0E-05 P08607	6.0E-05 P08607	8.0E-05 TS	6.0E-05 A	8 0F-05 R75639 4	6.0E-05 A	8.0E-05/A	6.0E-05 BE858403.1	5.0E-05 AW392086.1
	Expression Signal	5.72	1.71	0.95	0.71	1.09	2.97	3.09	1,69	1.69	1.19	1	1	3.07	3.61	3.61	1.4	0.95	0.78	0.76	0.62	2.22	0.63	1.21	1.21	0.85	0.59	308	3.36	14.34	1.4	10.48
	ORF SEQ ID NO:			29776	30221	33954	35175		27484	27485	28046	28137	28138	26080	31541	31642	32066	32520	33803	33804	34172	34177	34316	35050	35051	35271	35462	36579	37415	30822		26802
	Exon SEQ ID NO:	15919		1	. 1	- 1	- 1	23782	14733	14755	15309	15399	1		18611	18611	19081	19496	20678	20678	21035	21040	21171	21882	21882	22099	22274	23340	24103	25239	25053	14129
	Probe SEQ ID NO:	3156	4339	4412	4871	8124	9453	11112	2020	2020	2595	2690	2690	2817	5822	5822	6309	6834	7983	7983	8342	8347	8479	9151	9151	9421	9621	10649	11502	12387	12810	1382

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	Top Hit Descriptor		EST79996 Placenta I Homo sapiens cDNA similar to similar to p53-associated protein	EST79996 Placenta I Homo saplens cDNA similar to similar to p53-associated protein	9gment HS21C102	OFFINATION RECEPTOR-LIKE 1 (G-PROTEIN COUPLED RECEPTOR DEZ) 4084c10.x1 Soares_feta_liver_spleen_1NFLS_S1 Homo capiens cDNA close in an open country.	contains Alu repetitive element; contains element KER repetitive element;	AU125721 NT2RM4 Homo sapiens cDNA clone NT2RM4002075 5'	Mus musculus myosin light chain 2, precursor lymphocyte-specific (Mylc2al), mRNA	8-98	8 6-8	00105/451F1 NIH_MGC_21 Home sepiens cDNA clone IMAGE:3842292 5	Accounts of an age of the series of the series and the series of the ser	nieredo XI NCI, CGAP, Lu24 Homo sapiens oDNA clone IMAGE:3009838 3/	riomo sapieno interfeuidin-1 receptor antagonist homolog 1 (iL1HY1), mRNA	IGEN 8 (MAGE-8 ANTIGEN)	re(1)-acid glycoprotein gene	IV Homo sapiens cDNA 5' end	Mg300109.XT Sogres NSF FB 9W OT PA P S1 Homo sapiens cDNA clone IMAGE:2387209 31	PROTEIN KINDS CHEINDING PROTEIN NELL2 PRECURSOR (NEL-LIKE PROTEIN 2)	Homo serviers Diocesson and Service Nells Precursor (Nell-Like Protein 2)	qh8e11.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1855052.3' similar in contains		2448a12.rt Stratagene hNT neuron (#837233) Homo sanjans cDNA class (14A OF come of the com	contains Alu repetitive element, contains element L1 repetitive element:	1319 Homo sapiens cDNA	ne, complete ods	oviral like element	18 left arm of chromosome XV	DAR ZPOSIU64_T1 666 (synonym: hfkd2) Homo sapiens cDNA clone DKFZp6661064 6	OUTSOMBER I NINT MISC 44 Home saplens cDNA clone IMAGE:3608653 5' Home saplens TNNT1 gene, excris 1-11 (and lained CDS)
Single Exon Probes Expressed in Brain	Top Hit Database Source	1	7	7	T	T	T	Т	Mus musculus myosin light chain	Homo saplens SYBL1 gene, exons 6-8	Т	Т	T	T	T	T	7	Т	T	T	T	Т	7	T^{-}	T	T	Homo sapiens p47-phox (NCF1) gene, complete ods	H.sapiens DNA for endogenous retroviral like element	Т	Т	\top
ngle Exo	Top Datak Sour		ESI HUMAN	ESI HOMAN	SWISSDOOT		ESI HUMAN	ES HOMAN	Z P		N. I. I. I. I. I. I. I. I. I. I. I. I. I.	EST HOWAN	ENT LINEAN		CANTO COLO	SWISSPING NT	EST HIMAN	TANK IN THE	SWISSPECT	TOGGSTWS		Ton Land	NE TOWAR		EST_HUMAN	ESI HUMAN	2 2		NI Got Unitable	EST LIMAN	NT.
S	Top Hit Acess No.	3 OF OF A ASSOCIA	05 A A SOOOTS 1	3.0F-05 AI 169302.2	3.0E-05 P97468	A1240004 4	05 A1 (195721 1	44070400 11	A 12257	16 A.1225782 4	16 BF 733157 1	16 AA284049 1	3.0E-05 AW770982 1	6912434 NIT	12131	3.0E-05 X03273 1	-	T	T	T			2.0E-05 M13792.1		AA160562.1	T	T			T	П
	Most Similar (Top) Hit BLAST E Value	20 0 0	3 05 05	3050	3.0E-05	3.05.05	3.0F-08	30F-05	3.0F.05	3.0E-05	3.0E-05	3.0E-06	3.0E-06	3.0E-05	3.0E-05	3.0E-05	3.0E-06/	3.0E-06/	3.0E-05 Q62918	3.0E-05/C	3.0E-05 L77570.1	2.0E-05 A	2.0E-05 A		2.0E-05 A		2 05 05 X	2 OF -05 X	2.0E-05 A	2.0E-05 BI	2.0E-05 A.
	Expression Signal	760			-	0.82	0.97	1.66	1.17	1.17	2.33	1.47	1.58	123	0.51	0.58	1.22	292	0.98	0.98	1.77	1.09	2.43	7 / 7	1.23	0.93	122	0.71	0.78	1.09	1.92
	ORF SEQ ID NO:	29798	29799		30065	26082	30144	31161	32433	32434	33616	34082	34630	34633	34638		35068		36303	38304		27784	28037		28544	28754	28779	-			31382
	Exon SEQ ID NO:	17170	17170	17285	17420	13441	17522	18269	19419	19419	20494	20944	21483	21487	21491	21719	21899	22218	23079	23079	24585	15048	16301	15425	16899	16102	16121	16242	16539	17377	18467
	Probe SEQ ID NO:	4434	4834	4550	4686	4785	4791	6470	6999	6889	7799	8260	8791	8795	8799	9029	8220	9583	10433	10433	12072	2323	2687	2718	1		L	3485	L	4643	

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5827	18616		69:0	2.0E-05	-05 AF029308.1	۲N	Homo saplens chromosome 9 duplication of the T cell receptor beta locus and tryosinogen gene families
5880	18666	31606	0.76	2.0E-05	05 Q13183	SWISSPROT	RENAL SODIUMDICARBOXYLATE COTRANSPORTER (NA(+)DICARBOXYLATE COTRANSPORTER)
5880	18666	31607	0.76	2.0E-05	05 Q13183	SWISSPROT	RENAL SODIUMDICARBOXYLATE COTRANSPORTER (NA(+)DICARBOXYLATE COTRANSPORTER)
9065	18844	31808	0.61	2.0E-05	05 A1149272.1	EST HUMAN	qc72a02.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone IMAGE:1715114 3' similar to contains L1.t3 L1 repetitive element:
6527	19293	32297	2.26	2.0E-05	AA714330.1	EST HUMAN	nw06d12.s1 NCI_CGAP_SS1 Hamo saplens cDNA clone IMAGE:12385193'
6801	19462	32483	3.27	2.0E-05	2.0E-05 Y08926.1	N	P.falciparum mRNA for AARP1 protein, partial
6814	19475	32497	1.12	2.0E-05	05 A1492960.1	EST HUMAN	qz47b06.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2030003 3' similar to TR:002711 002711 PRO-POL-DUTPASE POLYPROTEIN
6824	19485		9.37	2.0E-05	05 Al991025.1	EST_HUMAN	wu35h07.x1 Soares_Dieckgraefe_cdon_NHCD Homo sapiens cDNA clone IMAGE:2622077 3'
7053	19744	32806	1.93	2.0E-05	05 AF224262.1	ΤN	Heterodontus francisci HoxA10 (HoxA10), HoxA9 (HoxA9), HoxA7 (HoxA7), HoxA6 (HoxA8), HoxA6 (HoxA4), HoxA4 (HoxA4), HoxA3 (HoxA3), HoxA2 (HoxA2), and HoxA1 (HoxA4) gense, complete cds
7053	19744	32807	1.93	2.0E-05	05 AF224262.1	Į.	Heterodontus francisci HoxA10 (HoxA10), HoxA9 (HoxA9), HoxA7 (HoxA7), HoxA6 (HoxA6), HoxA5 (HoxA4), HoxA4,
7267	19951		0.83	2.0E-05	05 AF128847.1	N	Homo sapiens indolethylamine N-methyltransferase (INMT) mRNA, INMT-2 allele, complete cds
7785	20480	33605	1.71	2.0E-05	05 AI381040.1	EST_HUMAN	tg20h05.x1 NCI_CGAP_CL1 Homo sapiens cDNA clone IMAGE;2109369 3'
9020	21710	34862	.0.53	2.0E-05	05 BE244840.1	EST_HUMAN	TCBAP2E1590 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project≔TCBA Homo sapiens cDNA clone TCBAP1590
90 20	21710	34863	0.53	2.0E-05	05 BE244840.1	EST HUMAN	TCBAP2E1690 Pediatric pre-B cell acute lymphoblastic leuksmia Baylor-HGSC project≔TCBA Homo saplens cDNA clone TCBAP1590
9167	21837	35002	0.58	2.0E-05	P49457	SWISSPROT	COMPLEMENT DECAY-ACCELERATING FACTOR (CD55)
9167	21837	35003	0.58	2.0E-06	2.0E-06 P49457	SWISSPROT	COMPLEMENT DECAY-ACCELERATING FACTOR (CD66)
9823	22474	35677	0.49	2.0E-05	١	- E	Homo sapiens chromosome 21 segment HS21C007
10035	22683	35900	0.87	2.0E-05		Т	7175g09.y1 NCI_CGAP_Bm20 Homo sapiens cDNA clone IMAGE:3340576 5
10457	23103	36333	0.54	2.0E-05	2.0E-05 AJ131024.1	TN	Homo sapiens otase gene, exon 1-alpha
10457	23103	36334	0.54	2.0E-05/	D5 AJ131024.1	TN	Homo sapiens otase gene, exon 1-alpha
10489	23135	36362	1.98	2.0E-05	05 N41751.1	EST_HUMAN	W91806.11 Soares_placenta_8to9weeks_2NbHP8to9W_Homo sapiens cDNA clone IMAGF:259570 57

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Table 4
Sindle Exon Probes Expressed

				ТТ	_	_			_	_	_	_				_				_		_	_				
Single Exon Probes Expressed in Brain	Top Hit Descriptor		Wested of Scares, placenta, 8tc9weeks, 2NbHP8tc9W Homo saplens cDNA clone IMAGE,289670 8'	130h09.X1 NOL CGAP Gas4 Homo sapiens cDNA clone IMAGE:2522077 3' 130h09.X1 NOL CGAP Gas4 Homo sapiens cDNA clone IMAGE:2132033 3' similar to TR:Q13538 Q13538	#30h09.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2132033 3' similar to TR-013838 013828	UNEZ: FUNCTION UNKNOWN.	hw21a03.x1 NCI_CGAP_KId11 Homo sapiens cDNA clone IMAGE:3183532 3' similar to TB:010822	WIZEZ GLYCOPHORIN HEPZ;	tioning septems. ABCA1 (ABCA1) gene, complete cds	Human gene for L-histidine decarboxylase, complete cds	POTATIVE SERINETHREONINE-PROTEIN KINASE P78	Homo saplens chromosome 21 segment HS21C082	Drosophila melanogaster strain Lamto 120 Suppressor of Hairless (Su(H)) gene, partial cds Homo sapiens calclum channel alpha1E subunit (CACNA1E) and	spliced	MOSAIC PROTEIN LGN	Homo sapiens chromosome 21 segment HS21C003	zw69g04.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:781494 5	xy49g11 x1 NCL CGAP_Lu34.1 Homo sapiens cDNA clone IMAGE:2856548 31	noting septents Spass gene for spassfin protein ns19g02.s1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:1184114 3' similar to containe 1.1.4.1.4	Li repaidive element ;	north Spring prospholipase Az, group X (PLA2G10) mRNA, and translated products	Homo senions chemical (SJOGREN SYNDROME TYPE A ANTIGEN (SS.A.)) (RO(SS.A.))	2336H2.s1 Seares directions 2 t segment HSZIC027 2336H2.s1 Seares directions NDZHES BW Homo capiens cDNA clone IMAGE:788519 3' similar to	2805611.11 NC_CGAP_GCB1 Homo septems cDNA clone IMAGE 684332 ff circles to contact at	repetitive element; contains element TAR1 repetitive element;	AV AZTBU FILL Flomo septiens cDNA clone HTFBIH01 5: hd41b0Zx1 Soares_NFL_T_GBC_S1 Homo septiens cDNA clone IMAGE;2912043 3; similar to contains	OFR.t1 OFR repetitive element;
gle Exon Pro	Top Hit Database Source	FOU	EST HUMAN	EST HUMAN	TOT	EST HUMAN	100	ES TOWAN	114	100000	7	2 12		\neg	/ISSPROT	7	Т	ESI HUMAN		NAMOL	/Icepport	Т	HEMAN	Т	EST HUMAN		EST HUMAN O
รัฐ 	Top Hit Acession No.	15 N41751 1	2.0E-05 AI991025.1	2.0E-05 AI493285.1	2.0E-05 A1403285 1			T	I	T	200	T	T	-	1	AL103203.2	T	T		5844	P19474	27.2			AV732400 4		7
	Most Similar (Top) Hit BLAST E Value	2.0E-08	2.0E-05	2.0E-05	2.0E-05	2.0E-05	2 0F-06	2.0E-05	2.0F-05	1.0F-05 P27448	1 OF OB	1.01-05			1.05-05	1.05.00	10505	1.0E-08	1 05-05	1.0E-05		1.0E-05 A	1.0E-05/A	40 HO		1 OF -05 A	
	Expression Signal	1.98	2.42	1.33	1,33	2.27	4.86	2.27	1.44	3.22	18	1.91		200	4.0	2 50	1 84	122	2.58	3.28	1.16	2.24	3.02	12.45	0.62	0.74	
	ORF SEQ ID NO:	36363		37243	37244	36428			ļ.	27731	28143	29034		20337	29523	29608	30152	32428	32530	32722		H	34799	35022	35189	35593	
	Exon SEQ ID NO:	23135	19485	23948		23197	25168	25155	26247	14991	15603	16394	16545	16690	16894	16985	17530	19415	19505	19675	20341	21505	21649	21857	22022	22389	
	Probe SEQ ID NO:	10489	10541	11287	11287	11430	12185	12342	12674	2265	2700	3641	3783	3949	4152	4244	4799	9883	6980	6982	7677	8813	8968	9187	9566	9738	

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	Top Hit Descriptor	hd41b02.xf Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE::2912043 3' similar to contains OFR:tl OFR repetitive element ;	UI-H-Biz-agk-a-08-0-UI.s1 NCI_CGAP_Sub4 Homo sapiens cDNA clone IMAGE:2724398 3	UI-H-BIZ-agk-a-08-0-UI.s1 NCI_CGAP_Sub4 Homo saplens cDNA clone IMAGE:2724398 3'	he07c10.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2873010 3' similar to contains L1.tZ L1 repetitive element;	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphale transporter (NPT3) gene, complete ods	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HIA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene	#73a08.x1 NCI_CGAP_HSC3 Homo sapiens cDNA clone IMAGE:2246386 3'	qg11b08.x1 Soares_placenta_8tc9weeks_2NbHP8tc9W Homo sapiens cDNA clone IMAGE:1759191 3'	Human elanine:glyακ/late aminotransferase (AGXT) gene, exons 1 and 2	Homo sapiens differentiation antigen CD20 gene, exons 5, 6	RC1-BT0313-110500-017-a07 BT0313 Homo sapiens cDNA	LINE-1 REVERSE TRANSORIPTASE HOMOLOG	ox20g01.x1 Scares_fetal_liver_spieen_1NFLS_S1 Homo sapiens dJNA cione INAGE:1636912 3 similiar to contains Alu repetitive element,	Homo sapiens chromosome 21 segment HS21C009	SUSHI REPEAT-CONTAINING PROTEIN SRPX PRECURSOR (DRS PROTEIN) (DOWN-REGULATED	BY VSRC)	SOSTI REPEAT-CONTAINING PROTEIN SON AT NECONSON (50.01 1.00 ELLY) (50.00 Human apolipoprotein E (APOE) gene, hepatic control region HCR-2	PUTATIVE SERINE/THREONINE-PROTEIN KINASE C22E12.14C	RC3-CT0283-201199-011-h11 CT0283 Homo sapiens cDNA	FASCICLIN II, PHOSPHATIDY (INOSTIOL-LINKED ISOFORM PRECURSOR (FAS II)	FASCICLIN II, PHOSPHA I IDYLINOSI I OL-LINKED ISOPOKM PRECORSOR (FAS II)	
	Top Hit Database Source	EST_HUMAN OF	EST_HUMAN UI-	EST_HUMAN UI	had EST_HUMAN rep	孔 孔 孔	II)	NT	EST_HUMAN #7	EST_HUMAN 49	NT Hu	NT Ho		SWISSPROT LIN	EST_HUMAN 00	NT Ho		SWISSPROT BY	SWISSPROT BY	NT	SWISSPROT PL	\neg		SWISSPROT F
	Top Hit Acession No.	1.0E-05 AW 510902.1		1.0E-05 AW 291521.1	1.0E-05 AW466995.1	1.0E-05 U91328.1	1.0E-05 U91328.1	1.0E-05 AF111167.2	9.0E-06 AI583811.1	9.0E-06 A 1218983.1	9.0E-06 M61755.1	9.0E-06 L23416.1	42.1	P08547	9.0E-06 A1034370.1	9.0E-06 AL163209.2		9.0E-06 Q63769	9.0E-06 Q63769	9.0E-06 U35114.1	9.0E-06 Q10384	8.0E-06 AW362539.1	8.0E-06 P34083	P34083
	Most Similar (Top) Hit BLAST E Value	1.0E-05	1.0E-05	1.0E-05	1.0E-05	1.0E-05	1.0E-05	1.0E-05	9.0E-06	9.0E-08	9.0E-06	9.0E-08	9.0E-06	9.0E-06 P08547	9.0E-08	9.0E-08		9.0E-06	9.0E-08	90E-06	9.0E-06	8.0E-06		8.0E-06 P34083
	Expression Signal	0.74	1.18	1.16	1.87	1.97	1.97	1.38	4.8	3.53	2.82	2.61	8.0	0.85	13.94	1:1		2.69	2.69	4.3	3.46	1.27	0.75	0.75
	ORF SEQ ID NO:	35594		35670		36760	36761	37780	28129	28498		31532	32534	33099	33450			34715	34716	١	ĺ	27986		36299
•	Exan SEQ ID NO:	22389	22467	22467	22732	23518	23518	24438	l	15857	16350	1	19509	20021	20337	1		21572	24572	_		Ĺ.,	23076	23076
	Probe SEQ ID NO:	9738	9816	9816	10084	10836	10836	11854	2678	3092	3597	5815	6765	7340	7673	8363		8881	8881	9122	10858	2532	10430	10430

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		3' similar to contains		ontains Alu repetitive				07	mRNA					3' ofmilar to							mplete cds						ains Alu	Alu	-
Single Exon Probes Expressed in Brain	Top Hit Descriptor	ab30f10.s1 Strategene lung (#837210) Homo sapiens cDNA clone (MAGE:854251 3' similar to contains MER20 t1 MER20 renealities closes and the contains about the contains of the contains and the contains and the contains and the contains are contained to contain the contains and the contains are contained to contain the contains and the contains are contained to contain the contains and the contains are contained to contain the contains and the contains are contained to contain the contains are contained to contain the	Homo sapiens KIAA0555 gene product (KIAA0555)	qw16g09.x1 NCI_CGAP_UI3 Home sapiens cDNA clone IMAGE:1991296 3' similar to contains Alu repetitive	BST99205 Throld Home confers about at	OV2-OT0062-250400 172 but OTAGE I	W65607.17 Soares_multiple_sclerosis_2NbHMSP Homo sanians_cDN4_close_instanc	THOUSE THE PROPERTY OF THE PRO	CERTIL OF ASAMN BEEN 1700 F. 1	SOURSESSES IN MOS 57 !!	CONTRACTOR DATASES AND THE SECOND CONTRACTOR OF THE SECOND	CV3-B10370 A40300 405 44 PTS 200 Homo sapiens cDNA	OVARIAN ARI INDANT MESSACE BEATEN (2015)	oxoboxx1 Social liver spleen_1NFLS_S1 Home sepiens cDNA clone IMAGE:1656738 3' similar to	Contains MEK8.t2 MEK8 repetitive element;	was nuscalus E-cadherin anding protein E7 mRNA, complete cds	11 5-1 MO070-140/A00 GES -AST 11/25=5:11	John Saplens calclum change 1 1 1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Homo sapiens chromosome 21 serment HS21Ch4e	iman ABI was asset to be a set of the set of	Homo saniers gaze for 1 ECT2 and littor 12, and putative M8604 Met protein (M8604 Met) gene, complete cds	SCA-CTO302-130200 043 PG2 CTO302 11	RC1-CT0302-120200-013-F02 CT0302 Home sapiens cDNA	EST185498 Colon carringma (UCC)	COMPI FMENT C2 BBECT IBSOB (22)25 CONTINUES CONA 5' end	HA0877 Human fetal liver cDNA library Homo contract CNA	948-03.71 Source Infant brain 1918 Home septens CDNA clone IMAGE:53254 5' similar to contains Alu	xc68g12x1 NCI_CGAP_Eso2 Homo septens cDNA clone IMAGE:25895743' similar to contains Alu	repetitive element contains element MER21 repetitive element;
igle Exon Pro	Top Hit Database Source	EST HUMAN	N	NAMI H FOR	EST HUMAN	EST HUMAN	EST_HUMAN	L	SWISSPROT	EST HIMAN	FST HIMAN	FST HIMAN	SWISSPROT	П	Т	SWISSPROT	T	Т				T HUMAN	Т	EST HUMAN	1	Т			ESI HUMAN IN
ris	Top Hit Acession No.	-08 AA669729.1	7682177 NT	-06 Al368252.1	-06 AA385542.1	-06 AW883141.1	7.0E-08 N98845.1	TN 60702411	7.0E-06 Q61147	Γ							T	157	5.0E-06 AL 163248,2		=			l	l		6 R16267.1 E		7
100	Most Similar (Top) Hit BLAST E Value	7.0E-06	7.0E-06	7.0E-06	7.0E-06	7.0E-06	7.0E-08	7.0E-08	7.0E-06	7.0E-06	8.0E-08	6.0E-08	8.0E-06	8.0E-06	6.0E-06	8.0E-08 Q02040	8.0E-08	8.0E-06	5.0E-06	5.0E-06 U07561.1	5.0E-06	5.0E-08 A	5.0E-08 A	5.0E-08 A	5.0E-06 P	5.0E-06 A1065045.1	4.0E-06 R	4 0F-08	יייייייי
	Expression Signal	2.69	3.42	5.83	0.92	5.68	1.01	0.7	0.45	2.32	1.28	1.08	1.91	2.21	1.32	1.08	1.48	2.27	3.27	2.31	1.1	0.53	0.63	6.16	0.45	2.83	6.1	70.7	
	ORF SEQ ID NO:		26851			1	31429	34524		30808	28329	29076	28359	30074	30700	30787		30979	31713	31983	32887	34192	34193	35862	36273	30987	26046	26266	
\vdash	SEQ ID NO:		14167		LI	J	18508	!		_			15708	17442	18071	18127	22407	25016	18752	19019	19821	21052	21062	22650	23056	24953	13411	13596	
	Probe SEQ ID NO:	958	1419	2876	3551	2609	6715	8688	88 8	11930	2918	3680	4705	4710	5285	5324	9226	12755	2970	6245	7134	8329	8359	10002	10410	12649	632	828	

Page 199 of 536 Table 4 Single Exon Probes Expressed in Brain

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		_	T	T	T	Т	Т	T	ē 7		7						_	_	_	_			_					_	
Single Exon Probes Expressed in Brain	Top Hit Descriptor		HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	KNOB-ASSOCIATED HISTIDINE-RICH PROTEIN PRECURSOR /KAHRE)	AV657555 GLC Homo saplens cDNA clone GLCFDB05 3'	2002e05.r1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA close IMA OF FACES 21	UI-H-BI3-aky-g-05-0-UI.s1 NCI_CGAP_Sub5 Homo saplens cDNA clone IMAGE 2724478 s	Mus musculus gene for odorant receptor A16, complete cds on34h01.s1 NCL CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1668809 3's similar to contains A1.	etanent; te51f06.x1 Soares NFL T GBC S1 Homo saniene cDNA alassa MAAAF assassa deed	Q13537 MER37 TRANSPOSABLE ELEMENT, COMPLETE CONSENSUS SEALITIONS	wj90b04.x1 NCI_CGAP_Lym12 Hamo saplens cDNA clane IMAGE-2446nes s	MR3-SN0067-120400-002-f02 SN0067 Homo sepiens cDNA	A447R Heart Homo saplens cDNA clone A447	zh27c11.s1 Soares, pinsel, gland_N3HPG Homo seplens cDNA clone IMAGE:413300 3' similar to	yu37c04,r1 Sogres overy tumor NbHOT Homo seplens cDNA close MACE conserver	KERATIN, TYPE II CYTOSKELETAL 8 (HUMAN);	Homo saplens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	Homo saplens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	11097U.X1 NCI_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2141730 3/	yw66e03.s1 Soares_placenta_8to9weeks_2NbHP8to9W Home emigra CNM_101_111111111111111111111111111111111	AV748969 NPC Homo saplens cDNA clone NPCAXD05 5'	PRO EIN MOV-10 ORGANIC CATION/CARNITINE TRANSPORTED 3 /601 HTF 01551	AFFINITY SODIUM-DEPENDENT CARNITINE COTBANISPORTER	Mus musculus D6MM5E protein (D6Mm5e) mRNA, complete cols	MEROZOITE SURFACE PROTEIN CMZ-8	Homo saplens chromosome 21 segment HS210078	208a12.s1 Sogres_feta_liver_spleen_1NFLS_S1 Homo sapiens cDNA olone IMAGE;429982 3' similar to contains Au repetitive element:	206a12.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:428882.31 elmiliar to	Ovariants Aur repetitive element; DNA-DIRECTED RNA POI VAIEBASE IIIT ABASEA AURTHUS
igle Exon Pro	Top Hit Database Source		SWISSPROT	SWISSPROT	ES! HUMAN	EST_HUMAN	ES HOMAN	I HOUSE	NAME OF THE PARTY	П	٦		EST_HUMAN	EST_HUMAN		HOMAN	Z		\neg	EST HUMAN	_		/ISSPROT		SWISSPROT	\neg	EST_HUMAN co	NAMI H TSE	Т
Sir	Top Hit Acess No.	DOVOZO	DOG 740	2.0E-08 AVAETEE 4	2 0E 08 A A 175 40 4	AM/450345.1	2.0E-08 ABARABBA 1			T	2 OF OR A MOSSOCO	T		AA772497.1	Labora 4]	T	T		-	AV /40909.1			Ar084364.1		AL1032/8,2 N	AA034141.1 ES	AA034141.1	
	Most Similar (Top) Hit BLAST E Value	2 OF OR	2.0E.	l			2.0E-06	2.0E-06	100	2.0E-00	2.0E-U0	2.05.00	20E-06	2.0E-06	200-08	2 0F-08 4	2.0E-08	2.0E-06		2.0E-06 N	20F-081		1.01-08	1.0E-06 A	10E-00 F		1.0E-06 A/	1.0E-06 AA	1.0E-06 P2
	Expression Signal	2.37	1.68	112	1 59	0.82	1.82	0.63	ca c	5.47	8	3 2	10.0	9.0	8	0.82	0.82	0.46		- E	2.1	8	2.30	1 81	1 67		1.27	1.27	1.34
	ORF SEQ ID NO:	27926			L		L		31768	32108		33809		1	34586	34969	34970		00720	92	30609	25477	28080	78888	26947	-	26997	26998	
	SEQ ID NO:	l i				16505	16510	18779	18808	L	ı	20683	ı	21427	21439	21804	21804	21823	22263	22470	25357	12860	13421	14181	14261	\vdash	14311	14311	14325
	Probe SEQ ID NO:	2469	2571	3508	3744	3763	3768	5998	8028	6348	7818	7988		8735	8747	9116	9116	9135	agor	9819	12251	32	842	1434	1514	-	1564	1564	1578

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Human ABL gene, exon 1b and intron 1b, and putative M8604 Met protein (M8604 Met) gene, complete cds qp54e02.x1 NCI_CGAP_Co8 Homo sapiens cDNA clone IMAGE:1926842 3' qv23f06.x1 NCI_CGAP_Lym6 Homo sapiens cDNA clone IMAGE:1982435 3' similar to contains element 2442002.s1 Stratagene hNT neuron (#837233) Homo sapiens cDNA clone IMAGE:632354 3' similar to zq42c02.s1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:632354 3' similar to zo04d11.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:785493 3' similar to za55e01 s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE 296472 3 zo17e08.r1 Stratagene colon (#837204) Homo sapiens cDNA clone IMAGE:587174 5' ol29c08.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone (MAGE:1524878.3) Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions Homo sapiens glypican 3 (GPC3) gens, partial cds and flanking repeat regions Homo sapiens chromosome 21 segment HS21C080 gb:D26129 RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN); Homo sapiens shox gene, alternatively spliced products, complete cds Homo sapiens shox gene, alternatively spliced products, complete cds Top Hit Descriptor CM0-BT0281-031199-087-h04 BT0281 Hamo sapiens cDNA FIBRINOGEN ALPHAALPHA-E CHAIN PRECURSOR MR3-FN0004-090600-001-e04 FN0004 Homo saplens cDNA RC4-NT0054-120500-012-b03 NT0054 Homo sapiens cDNA MR1-BT0800-030700-002-c06 BT0800 Homo sapiens cDNA MR3-FN0004-090600-001-e04 FN0004 Hamo sapiens cDNA L5-HT0730-020500-074-g01 HT0730 Homo sapiens cDNA DYNEIN GAMMA CHAIN, FLAGELLAR OUTER ARM Homo sapiens p47-phox (NCF1) gene, complete cds Homo sapiens p47-phox (NCF1) gene, complete cds Homo sapiens chromosome 21 segment HS21C003 Homo saplens p47-phox (NCF1) gene, complete cds Homo sapiens p47-phox (NCF1) gene, complete cds Homo sapiens chromosome 21 segment HS21C081 SW:POL_SMSAV P03359 POL POLYPROTEIN; SW:POL_SMSAV P03359 POL POLYPROTEIN 15 KDA SELENOPROTEIN PRECURSOR Single Exon Probes Expressed in Brain MIR repetitive element EST HUMAN EST HUMAN EST HUMAN SWISSPROT EST_HUMAN EST_HUMAN EST HUMAN SWISSPROT **EST HUMAN** EST_HUMAN EST_HUMAN **EST HUMAN** EST_HUMAN Top Hit Database Source EST_HUMAN EST HUMAN EST_HUMAN SWISSPROT Ä 눋 z 뉟 눌 Top Hit Acession 1.0E-06 BE834518.1 1.0E-06 BE834518.1 1.0E-06 O60613 1.0E-06 BE063527.1 1.0E-06 P02671 1.0E-06 AW 890941.1 9.0E-07 AF003529.1 9.0E-07 AL163280.2 1.0E-06 AF184614.1 1.0E-06 AF184614.1 1.0E-06 AA449257.1 1.0E-06 AA164914.1 1.0E-06 AA164914.1 9.0E-07 AF003529.1 BF333015.1 1.0E-06 AA132611.1 1.0E-06 AF184614.1 1.0E-06 AF184614.1 9.0E-07 AL 163281.2 1.0E-06 BE185330.1 1.0E-06 AA912623.1 ġ 1.0E-06 U07561.1 1.0E-06 BF333015 1.0E-06|AI347010.1 1.0E-06 AI287878.1 1.0E-06 N74635.1 1.0E-06 Q39575 1.0E-06 .0E-06 (Top) Hit BLAST E Most Simila Value 5.09 5.09 0.93 1.22 8.73 0.93 6.91 1.31 0.55 4.76 3.37 1.68 3.85 1.38 1.38 2.24 0.53 2.87 Expression Signal 27443 29703 30638 27443 27444 25790 ORF SEQ 30665 34215 35255 35205 35417 35418 35470 37641 37642 37161 30666 33997 ΩÑ SEQ ID 18016 14723 24318 14723 22340 18189 19517 21080 22033 23031 24245 24318 14723 13150 13150 23875 Probe SEQ ID 4336 5232 5232 5389 5706 6773 7800 17900 171 9279 9581 9581 9627 11724 11724 351 11212 1987 1987 8387 9204 9688 11648 10385 12390

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Single Exon Probes Expressed in Brain	_
Top Hit Database Source	CM-BT178-220499-014 BT178 Hamo seplens cDNA
	EST_HUMAN
(Top) Hit BLASTE Value BLASTE Value B.0E-07 AF0879-8.0E-07 A128859 8.0E-07 A7128859 8.0E-07 A7128859 8.0E-07 A712834 8.0E-07 A7183381 6.0E-07 A7383981 5.0E-07 A1383881	5.0E-07 Al908587.1
Expression Signal 3.28 3.28 3.28 9.73 6.59 1.52 1.83 1.52 6.59 0.91 0.77 1.69 1.69 1.69 1.70 0.74 0.74 0.74 0.74 0.74 0.74 0.74 0	4.94
	36123
	22873
Probe SEO ID NO: NO: 11733 11733 1719 4719 4719 4719 4719 4719 6729 6964 6964 6964 6964 6968 8388	10265

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Single Exoli Flobes Explessed in Diam	Top Hit Descriptor	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	COLLAGEN ALPHA 1(1) CHAIN PRECURSOR	Homo sapiens Xq pseudoautosomal region; segment 1/2	QV0-CT0383-210400-204-612 CT0383 Homo sapiens cDNA	ws84h05.x1 NCL CGAP_Co3 Hamo sapiens CDNA clone IMAGE: 2004097 3	Homo sapiens SPP2 gene for secreted phosphoprotein 24 precursor, exams 1-o	HISTONE DEACETYLASE 8 (HD2) (HISTONE DEACETTESE WHDA1)	STONE DEACET YEARS (MDS) (MISTONE DEACET TEACH WILLDAY)	Homo sapiens chromosome z I segiment nozi coo?	xy49g11.x1 NO_CGAP_Lu34.1 Home sapiens clone invace.cocotto	601676748F1 NIH MGC 21 Homo sapiens clonk clone liwk GE 355551 5	601676748F1 NIH_MGC_Z1 Homo sapiens cunA cione ilwAcE:3939031 3	Homo saplens chromosome 21 segment HSZ1C018	WI81b08.x1 NCI_CGAP_KId12 Homo sapiens GUNA cione IMAGE. 2399703 3	wi81b08.x1 NCI_CGAP_Kid12 Homo sapiens cUNA clone IMA CETZ389703 3	PM1-BN0083-030300-003-e12 BN0083 Homo sapiens cDNA	Human microfibril-associated glycoprotein (MFAP2) gene, putative promoter region and alternatively spliced untansiated evens.	11e-remines Various ideas the company of the compan	normo sapiens Ad pseudosadosadira region, organismos	Interpretation of the property of the polytrophile interpretation of the polytrophile	Human igk subgroup i germine gene, excris i and 2, v-region ore and a containe Alli repolitive	nis8809,s1 NCL_CGAP_OVZ homo sapiens curva cione livrace. 300025 similar to contains and reposition element; contains L1, (3 L1 repetitive element;	Нитап polymorphic microsatellite DNA	MR0-BN0115-020300-001-f11 BN0115 Homo sapiens cDNA	MR0-BN0115-020300-001-f11 BN0115 Homo sapiens cDNA	yd50f12.r1 Soares fetal liver spieen 1NFLS Homo sapisns cDNA clone IMAGE:111695 6	HYPOTHETICAL 63,8 KD PROTEIN IN GUT1-RIM1 INTERGENIC REGION PRECURSOR	AV650201 GLC Homo sapiens cDNA clone GLCCCD01 3'	we86b12.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:234/961 3	yc14h09.s1 Stratagene lung (#837210) Homo sapiens cDNA cione (MAGE:80705 3' similar to similar to gb:M62982 ARACHIDONATE 12-LIPOXYGENASE (HUMAN)	yc14h09.s1 Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:80705 3' similar to similar to gb:M82982 ARACHIDONATE 12-LIPOXYGENASE (HUMAN)
le Exoli Piones	Top Hit Database Source	П	ISSPROT		П	T_HUMAN	1	┱	SWISSPROT HI	П	П		EST_HUMAN 60	_1	EST_HUMAN W							F F	EST HUMAN e	Г	EST HUMAN N	Г	Г	Г	EST_HUMAN A	EST_HUMAN W	EST_HUMAN g	EST_HUMAN 9
Buis	Top Hit Acession No.	7 P08547	17 P11087			1	5.1			١	07 AW419134.1	07 BE901975.1	١	07 AL163218.2	07 AI765528.1	07 AI765528.1	07 BE001828.1		0/ 0/8/ 18.1	3.0E-07 AJ271735.1	M99149.1	M64857.1	07 44526763 1	3.0E-07 M99149.1	3.0E-07 BE005077.1	BE005077.1	3.0E-07 T84704.1	07 P38739	07 AV650201.1	07 AI797236.1	-07 T57850.1	-07 T57850.1
	Most Similar (Top) Hit BLAST E Value	5.0E-07	5.0E-07	5.0E-07	5.0E-07	4.0E-07	4.0E-07	4.0E-07	4.0E-07 Q9Z2V6	4.0E-07	4.0E-07	4.0E-07	4.0E-07	4.0E-07	4.0E-07	4.0E-07	4.0E-07		3.05-07	3.0E-07	3.0E-07	3.0E-07	2 OF -07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07
	Expression Signal	1,28	4.04	2.52	3.48	2.02	0.83	1.74	1.74	9.0	5.41	0.47	0.47	0.49	3.14	3.14	1.66		9.64	2.12	2.67	2.03	1 40	1 83	7.61	7.61	1.16	1.45	7.42	0.86	1.3	1.3
	ORF SEQ ID NO:	36493	37413	-		29364		32923	32924	33644	34787	35890						_			26776			27749				L				
	Exon SEQ ID NO:	23256	24101	24173	25211	1	1	19854	J	l)	L	22675	22871	23536	1	1	1_			14101	14369	1,173	L	1	1	Į	1.	L		L	1
	Probe SEQ ID NO:	10560	11500	11574	12561	3981	7078	7167	7167	7823	8949	10027	10027	10223	10856	10856	11184		431	269	1353	1622	8	8000	2472	2472	3034	34.57	48.78	4711	5004	5004

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Table 4

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Probe SEQ ID NO:	Ban SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5059	17778	30396	0.84	2.0E-07	07 AW070995.1	EST_HUMAN	xa05h07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA Gore IMAGE.2007450 Similar to 17. 3000 E.C. CONTROL CE00923 PROBABLE RABGAP DOMAINS;
5260	1	30694	1.21	2.0E-07	07 AW898066.1	EST HUMAN	RCS-NNUCCOC-200405221-91 INVOCOTION September CDNA clone IMAGE:2734008 3'
6456	1			2.0E-07	07 AW448968.1	EST HUMAN	UL-TI-BIO-BRO-D-01-0-01-1 NOT Home sapiens CDNA clone IMAGE:1839177 3
9299	1	32337		2.0E-07	2.0E-07 AI208715.1	EST HUMAN	dgbodub,X1 Sustains again of the first of th
7568	ı		29.0	2.0E-07	07 X95159.1	LN	III. Septembrilla Direct your o contents of DNA plans HTCAEG02 5
8369	1			2.0E-07	07/AV729390.1	EST HUMAN	AV / 2839U TTC nome septems construction of the septems constitution of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the septems of the september of the septemb
8595	21287	34426	0.97	2.0E-07	2.0E-07 AA035198.1	ES! HUMAN	2xt you's 1 out of your or 1 series HS21C103
986	1		2.8	2.0E-	2.0E-07 AL163303.2	LN	Horito Saptem Cutoring 21 agriculture Saptem
10167	1_	36033		2.0E-	07 AW892507.1	EST HUMAN	CM4-NNUUUS-280300-124-830 1440300 15910 CAPETASE) (PROPERDIN FACTOR B)
900			60	2.0E	07 P00751	SWISSPROT	COMPLEMENT FACTOR B PRECONSON (SOC OF SOC OF
10300	_					1000	COMPLEMENT FACTOR B PRECURSOR (C3/C3 CONVERTABLE) (FROFENDIN) FOR CO. 2/C/C3 YOUR PICH RETA Q1 YOOPROTEIN) (GBG) (PBF2)
10386	3 23032	36246		2.0E	07 P00751	SWISSPRO	OET CHAPACAL 2017 CLT 2017 CLT 2017 HT0339 Homo sapiens cDNA
11871	1 24945	2	. 2.44	2.0E	-07 BE153717.1	EST HOMEN	The state of State of
11063	35242		2.39	2.05	-07 AI732462.1	EST_HUMAN	Ontains THR.b2 THR repetitive element:
1080	L	1 6	1.97	1.0E	-07 AL163282.2	LN.	Homo sapiens chromosome 21 segment nozi cook
300	L	L	1.11	1.8 B.1	-07 P10263	SWISSPROT	RETROVIAGOS-KELATED GAG TOLITICALET (TELECONET)
3 8	L	28945		1.0E	-07 P09256	SWISSPROT	GLYCOPROTEIN GPV
2000	L			1.0	-07 AL163282.2	NT	Homo sapiens chromosome 21 segment 1732 1 cook
17/6	L			1.06	1.0E-07 AV718662.1	EST_HUMAN	AV718662 GLC Homo sapiens cun ache Glucini off 3
4280	_L	29632		L	1.0E-07 AV718662.1	EST_HUMAN	AV718662 GLC Homo sapiens culva done occurred to
4690	1				1.0E-07 075820	SWISSPROT	ZINC FINGER FRO LEIN 100
5072	L	30406			1.0E-07 AA019181.1	EST_HUMAN	Zeoguzri Sua es reuna 1224 in transmissione family A2a (MAGEA2A), melanoma antigen family A12
	_						Homo seperas orruntiosomes was managen family A3 (MAGEA3), celtractin (MAGEA12), melanoma antigen family A3 (MAGEA3), celtractin (MAGEA12), melanoma antigen family A3 (MAGEA28), melanoma antigen family A3 (MAGEA28), celtractin (MAGEA12), melanoma antigen family A3 (MAGEA3), celtractin (MAGEA12), melanoma antigen family A3 (MAGEA13), celtractin (MAGEA12), melanoma antigen family A3 (MAGEA13), celtractin (MAGEA13), celtrac
6410	0 19178	32177	7 0.87		1.0E-07 U82671.2	LN	(CALT) NAD(P)H denydrogenase-like protein (NSO-110), and Errord (NSO-110), and Errord (NSO-110), and Errord (NSO-110) (NSO-110
6768	١				1.0E-07 BE047871.1	EST HUMAN	E43000.yi NO. COAF BINES Homo sepiens cDNA clone IMAGE:2291339 5'
6768	38 19512	32538			1.0E-07 BE047871.1	EST HUMAN	Expanso, J No. Contraction 100 September 200
7392	١				1.0E-07 N55081.1	EST HUMAN	IDNA TNIANZA ASIAKA AND TO THE TRIBOLA HOMO Sapiens CDNA
7548	<u> </u>	33320			1.0E-07 BF375909.1	ESI HUMAN	DAMA TRIONSA DROPE DOS TROOS Homo sapiens cDNA
7548	l				1.0E-07 BF375909.1	ES HOMAN	Home sapiens chromosome 21 segment HS210081
75/		46 33351	1.31		1.0E-07/AL163281.2	N	

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	Top Hit Descriptor			3		lomo sapiens cDNA clone IMAGE 434346 21	MOTIFS 8) (ADAMTS-8) (ADAM-TS8) (METH-2)	hu28h06.x1 NCI_CGAP_Mel15 Homo saplens cDNA clone IMAGE:3171416.31	MER 18	lane IMAGE:4274426 5'		ht53c11.x1 NCL_CGAP_Kid11 Home seriess - DNA	GIONE IMAGE:3132212 3' SIMILAR to TR:095722 CORT	1000	dex (germline)	oDNA clone IMAGE:2090195 3'	wn30a07.x1 NCI_CGAP_Gas4 Homo saplens cDNA clans MACE 221222	MINIOGE: 2440832 3' similar to contains OFR.12			oDNA clone IMA OF: 222222	le IMAGE:3943976 6'	e IMAGE:3943976 6'	cn15c02.x1 Normal Human Trabecular Bone Cells Homo sabiens CNA Acro Mutto	Control of the lacture of the lactur	no saplens cDNA clone NHTBC cn15c02 rendem	s cDNA		cros gene, complete cds; and unknown gene	garie, complete cds		
Origin Exon Probes Expressed in Brain			Г				\neg		Т	Г	Т	Γ				П		7	Homo sapiens chromosome 21 segment HS21C101	Ţ	Т	Т	1	T		N EST382776 MAGE resemigance, MAGE, and Cells Homo saplens cDNA clone NHTBC cn15c02 random	· -	Homo saplens jun dimerization protein gene partiel cd	٦	٦	٦	
SY SY			L _N	SWISSPROT	SWISSPROT	EST_HUMAN	SWISSPROT	EST HUMAN	EST HIMAN	EST HUMAN	Ā	1	ES HOMAN	Į.	EST_HUMAN	EST HUMAN	TOTAL TOTAL	TIV	Į.	TO TO	EST HIMAN	EST HUMAN		ES HUMAN	EST HUMAN	EST_HUMAN				SW ISSPROT		SWISSPROT
	Top Hit Acession No.	7 (000)	E-07 AL163203.2	1.0E-07 P97435	1.0E-07/P97435	1.0E-U/ AA683576.1	-07 P57110	-07 BE327843.1	Γ		1.0E-07 AL163282.2	-07 BE048770 4	7			S.UE-06/AV/34819.1	08/AI891052.1	Ī	Γ	Ī	T		8.0E-08 A1752387 4	T	8.0E-08 AI752367.1 ES	П	A E44467 0		I		\int	7
Most Similar	(Top) Hit BLAST E Value	100	200.	212	0 10	0-10-1	1.0E-07	1.0E-07	1.0E-07	1.0E-07	1.0E-07	1.0E-07	1 OF 07	00 00	00 00	8.0E-08	9.0E-08	9.0E-08	9.0E-08/	8.0E-08	8.0E-08	8.0E-08	8.0E-08		8.0E-08 A	8.0E-08 A	8 0E-08		7.0E-08/07	7.0E-08/X0	7.0E-08 P1	
	Expression Signal	0 64				200	À.	0.45	2:77	1.24	1.28	3.83	1.87	0.84	1.88		1.71	2.8	4.44	3.7	0.72	1.53	3.05		3.05	2.83	0.47		L	L	L	
	ORF SEQ ID NO:	33489	33941	33942	34691	35005		35353	80000	08000		30810		32940	35845		37082	3/587	1	1	+	1	34474	24475	35364		36338		25542	26764	28965	
	SEQ ID	20385	20808		1	21840	乚		L	23009	ı	25188	24864	19867	22438		243804	335	155.48 155.48	2000	18780		21330	21330	22180	-	23107	23874	12904	14088	16318	
Probe	SEQ ID NO:	7724	8114	8114	8853	9170	0.64	9836	9844	10362		12212	12514	78	9787	60,44	1168	12188	200	1020	3532		8638		9527	L	_4		_[$_{\perp}$	3563	

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	Top Hit Descriptor	DYNEIN HEAVY CHAIN (DYHC)	cong3.P11.A5 conorm Homo sapiens cDNA 3	Rattus novegicus Munc13-1 mRNA, complete cds	DYNEIN HEAVY CHAIN (DYHC)	DYNEIN HEAVY CHAIN (DYHC)	Homo saplens SCL gene locus	Home seriens chromosome 21 segment HS21 C048	Lano conjene chromosoma 21 segment HS21C048	Trusting services of the property of the prope	MINO-TILLOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCO	omo saprans night in the comment through the c	HOMB sapiests and analysis of any sapiests and sapiests and any sapiests a	NE-1 KEVERSE I MANSURIF I ASE ITOMOTION I MANOE 1102558 2 similar to contains	obsects, st NCI_CGAP_GCB1 Home sapiens cunA cione invAGE. 1555500 5 Sinital to commit	MER 12. b3 MER 12 repetitive element;	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: NEVENSE TRANSCONT 1902)	ENDONOCELASE)	Long capiens chromosome 21 segment HS21C103	AND COAD THA Hamp saplens cDNA clone IMAGE:943193 similar to contains Alu repetitive	element;	COMPLEMENT C2 PRECURSOR (CSICS CONVENTION)	QV0-CIUZZS-131U89-USH 1Z CIUZZS TUNIO SERIENIS CENTINA	DOROAL-VENTINAL PATTERNING TO LOND PROTEIN PRECURSOR	DOKSAL-VEN I NAL FAT LEXINO 1 CENTRAL STATEMENT CON CIONE DKFZp434J0426 5	UNITED ALCONOMINE AND AND AND AND AND AND AND AND AND AND	HOMO Saperis snot gene, and named spicos, process, contracting process, contracting and see (I IDRDASE)	URIDINE PHOSPHORICAGE (UDIVIDE)	J KANSMEMBRANE TROU CASE, OEL ME E	Croediles ginsels ribosofinal deficiency (Co. 2) in the contract ribosof	LINE-1 REVENUE I PANASCANI I PAGE 1 COMPANA CON Clone IMAGE: 1622903 3'	ot/odities Ocean Wilms immer Homo saplens cDNA clone IMAGE:1699411 3' similar to contains Alu	arized to A. Coosea Villes target ment of the separative element; repetitive element contains element MER22 repetitive element;
Single Evolutional Significance	Top Hit Database Source	SWISSPROT		12	Т	Т	Т				HOMAN		1	SWISSPROT		EST_HUMAN		ISSPROI		Ž		\neg		1	Т	EST HUMAN	Ł	SWISSPROT	SWISSPROT	NT NT		EST_HUMAN	EST_HUMAN
Billo Billo	Top Hit Acession No.		3.1								6.0E-08 BE144398.1	7662473 NT	6.0E-08 AL163248.2	P08547		6.0E-08 AA827075.1		P11369	6.0E-08 AL 163209:2	08 AL163303.2	08 AA493851.1	08 P06681	08 AW851878.1	08 P25723	-08 P25723	AL079581.1	4.0E-08 U82668.1	4.0E-08 P52624	015393	-08 L42571.1	-08 P08547	-08 A1016342.1	-08 AI050027.1
}	Most Similar (Top) Hit BLAST E Value	7.0E-08 P15305	7 OF 08 /	7 OF 08 1124070 1	7 OF 08	7.05-101-101-1	7.0E-081	7.0E-08/	6.0E-08	6.0E-08	6.0E-08	6.0E-08	6.0E-08	6.0E-08		6.0E-08		6.0E-08 P11369	6.0E-08	5.0E-08	5.0E-08	5.0E-08	5.0E-08	4.0E-08	4.0E-08	4.0E-08			4.0E-08	4.0E	4.0E-08	4.0E	4.0E
	Expression Signal	1 15	187	1.0	9.10	2.30	2.98	1.89	2.88	2.88	2.97	0.81	86'0	0.69		0.56		2.24	1.33	3.72	1.82	8.36	2.54	0.97		1.09	1.04	1.08	0.63	1.05	0.71	0.68	3.87
	ORF SEQ ID NO:	28086	20003		١		28966				27824		29588					37299		25547	7697	L	31085		27196		29284			l		<u></u>	2 36147
	Exon SEQ ID NO:	48340	1	- 1	-1	- 1		24975	13570	13570	l	15824	L	L	L	24006	L	23997		12909	14957	L	1	1	L	15655	16644	<u></u>	L	1_	L	L	4 22932
	Probe SEQ ID NO:	0000	202	10/16	11672	12619	12619	12690	798	798	2363	3058	4222	7044	3	7000	277	11391	11520	83	2220	11914	12095	1754	1754	2888	3894	8	(698	9037	9545	10228	10284

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	Top Hit Descriptor	2/76b08.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728247 5' similar to TR:G505579 G505579 NA/CA K-EXCHANGED .	278508.11 Soeres Lests NHT Homo saplens cDNA clone IMAGE:728247 5' similar to TR:G505579	INGER.;	2 Homo saplens cDNA clone IMAGE:4333300 5'	ZEG5g03.r1 Sogressian heart_NbHH19W Homo sepiens cDNA clone IMAGE:345556 5' similar to containo	t695611 x1 NCI_CGAP_C016 Home sapiens cDNA clone IMAGE:2062076 3' similar to contains MER18.b3	borgano in Line depours element; borgano in III MGC_10 Homo sapiens cDNA clone IMAGE:3048570 5' similar to TR:092158 092158		38 Homo sapiens cDNA clone IMAGE:1944045 5	21 segment HS21C046	th93h09x1 Soares NSF_F8_9W_OT_PA_P_S1 Homo sepiens cDNA clone IMAGE:2126273 3' similar to	TRANSPOSABLE ELEMENT, COMPLETE CONSENSUS SEQUENCE:	region	This law, at locates Int. 1 GBC, St Homo saplens cDNA clone IMAGE:1846294 31 yr12b10.st Soares breast Shall Homo saplens cDNA clone IMAGE:187195 31 similar to gb:M34079 TAT BINDING PROTFINA 1 (H) MANN.	7912910.81 Scares breast 3NHBst Homo sepiens cDNA clone IMAGE:187195 3' similar to gb:M34079 TATE BINDING PROTEIN 1 (HIMAN).	vg02f04.r1 Scares Infant brain 1NIB Homo saplens cDNA clone IMAGE:30948 5' similar to contains Alu	Home conjust the A	ZW48f07.r1 Soares, total_fetus_Nb2HF8 by Homo sepiens cDNA clone IMAGE;773317 3' similar to contains	S GENERAL MERKING REPORTING BETTEN TO	Deciz) mrNA, complete cds	39 OTODOS II	Home Colored Home Sapiens CLINA	Franco Sapiens CUNA clone IMAGE:3138883 5'	601570463F1 NIH MGC 21 Home earloan all 11 of the last
Single Exon Probes Expressed in Brain	Top Hit Descrip	276b08.1 Sogres_testis_NHT Homo sepiens cDNA clone IN G505579 NA/CA K-FXCHANGED	76b08.r1 Soares tests NHT Homo sapiens cDNA clone IN	6022480245 MACA,K-EXCHANGER.;	602248024F1 NIH_MGC 62 Homo sapiens cDNA clone IMAGE:4333300 5	2055903.r1 Soares fetal heart NbHH19W Home sapiens cDL1.r1 L1 repetitive element	Ф95а11 x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMA	TO TO THE MINE OF THE MAN SERVICE SERVICES OF THE MACE	STATAMIN 17.;	Harring No. 1 CAR Frze Homo sapiens cDNA clone IMAGE:1944045 5	nono sepiens chromosome 21 segment HS21C046	3h09x1 Soares NSF F8 9W OT PA P S1 Homo sapler	C1353/ C1353/ MER3/ TRANSPOSABLE ELEMENT, C	Short of grant MITC class 1 region	iquaturu, xi ocares IntGBC_S1 Homo saplans cDNA clone IMAGE:1845294 37 pr12b10.s1 Secrets breast 3NHBst Homo saplans cDNA clone IMAGE:187185 3' sin BINDING PROTEIN.1 (11) MANN.	yp12b10.s1 Soares breast by NbHBst Homo sapiens cDNA clon-BINDING PROTEIN-1 (HLIMAN).	vg02f04.r1 Sceres infant brain 1NIB Homo saplens cDNA clone	x8706x1 NCI CGAP 1.126 Homo contact DNIA 1	8f07.r1 Scarce, total fetus, Nb2HF8 9w Homo sapiens cD	Gallus dallus Dasha materi / Dasha Mark i a repetitive element	TOTORO 240200 684 - 00 OTTORO 24020 684 - 00 OTTORO 240200 684 - 00 OTTORO 240200 684 - 00 OTTORO 240200 684 - 00 OTTORO 240200 684 - 00 OTTORO 240200 684 - 00 OTTORO 240200 684 - 00 OTTORO 2402	MR0-OT0080-240200-01-908 OT0080 Homo sapiens cDNA	60115532151 NIH MCC 21 Home control and an analysis	Homo sapiens chromosome 21 segment HS24C047	TOURSE AND MOO OF IT
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	Expression Signal	17.1	1.71	4.02	4.02	4.31	2.18	2.22	4.24	1.66		3.56	0.52	1.32	61.58	61.58	2.27	9.03	9.14	1.01	13.62	13.62	24.4	2.38	12.18
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	Exon SEQ ID NO:	23681		23702	23702	25334	24887	18321	17955	20116		20313	22449	23626	24165	24165	24459	13014	13033	13269	13424	13424	13735	14069	14476
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Table 4
Single Exon Probes Expressed in Brain

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	TRICARBOXYLATE TRANSPORT PROTEIN PRECURSOR (CITRATE TRANSPORT PROTEIN) (CTP) (TRICARBOXYLATE CARRIER PROTEIN)	BONE MORPHOGENETIC PROTEIN 4 PRECITES OF ARIA 3	Homo sepiens major histocompatibility focis along ill conjust	Human lembda-imminocilobulin constant recipion	Homo sabians chromosome 21 seament HS21/Cn70	Homo sepiens chromosome 21 common US34 0570	ye58a12.51 Soares fetal liver spiego 1NFI S Home employed all and 11/2012.	quB6o11.x1 NOI_CGAP_Gas4 Homo septens cDNA done IMAGE:1978964 3' similar to contains L1.ts L1	9442607.x1 Soares fetal, heart, NbHH19W Homo saplens cDNA clone IMAGE:1732164 3' elmilar to	CMO-NN 1004-100300-273- of NN 1004-1	op74008.61 Soares NF. T. GBC S1 Home cantens child.	demonstrate DNA for a plant of the second of	7145e10.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sepiens cDNA clone IMAGE:3524443.3' similar to	contains MER29.b2 MER29 repetitive element;	Zoucuo,n Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE: 681992 6' similar to contains L1,12 L1 repetitive element:	Human lysosomal membrane glycoprotein-2 (1 AMP2) gene E' and and familing seed.	60111173F1 NIH MGC 16 Homo saplens oDNA clane IMAGE 335434 F	## ## ## ## ## ## ## ## ## ## ## ## ##	ve58a12.s1 Soares fetal liver splean 1NFI S Homo company CDNA class 144 OF 2002000	DKF20434C0514 rt 434 (singman: https://dx.com/com/com/com/com/com/com/com/com/com/	PM1-HT0527-160200-001-h05 HT0527 Home salvens convA digne UNF2p434C0614 6	XN85h08.X1 Spares NF T GBC S1 Home septem of NA Alexandra S1	MR3-HT0446-260300-201-h12 HT0448 Home seniors - DNA	OTIO Sablens (ihrohiset arough fodes seconds 2 / 1	Homo saplens testis-specific kinasa suhshtota (TeKS) man	745e10.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sepiens oDNA clone IMAGE:3524443 3' similar to contains MER29.b2 MER29 repetitive element;	
gle Exon Prok	Top Hit Database Source	SWISSPROT	SWISSPROT	LN L	L'A	NT	N	EST HUMAN			Т	Т	Į.		ESI HUMAN	EST HUMAN	Т	EST_HUMAN	EST HUMAN	Т	T	Т	Т	Т			EST_HUMAN o	
Sir	Top Hit Acession No.	08 P79110	-08 P98063	-08 AF044083.1	1.0E-08 X51755.1	9.0E-09 AL163279.2	9.0E-09 AL163279.2	9.0E-09 T97950.1	09 AI270615.1	09 A1183500 1	-		09 D86842.1		7.0E-08/DF-108/39.1	7.0E-09 AA256200.1	7.0E-09 L09709.1		7.0E-09 AA058626.1		6.0E-09 AL040439.1			6.0E-09 BE161653.1	4503710 NT	6.0E-09 AF200923.2		
	Most Similar (Top) Hit BLAST E Value	1.0E-08	1.0E-08	1.0E-08	1.0E-08	9.0E-09	9.0E-09	9.0E-09	8.0E-09	8.05-09	8.0E-09/	8.0E-09	7.0E-09	100	BO-UO.	7.0E-09	7.0E-09 L	7.0E-09 E	7.0E-09 A	7.0E-09 T97950.1	6.0E-09	6.0E-09	8.0E-09	6.0E-09	6.0E-09	6.0E-09 A	6.0E-09 BF108755.1	
	Expression Signal	1.2	0.77	4.14	2.82	4.65	4.65	0.52	0.62	7.66	2.65	2.65	1.73	å	10.0	0.82	2.91	1.42	0.6	1.49	0.99	3.12	11.59	0.93	1.96	3.76	1.44	
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	Exon SEQ ID NO:	. i			. 1	- 1	16959	22610	19159	19861	20594	21678	16346	20497		20641	21829	22731	22892	23266	14879	17650	18101	21167	21763	22824	23324	
	Probe SEQ ID NO:	9866	10453	11285	12282	4218	4218	8862	6390	7164	7899	8887	3593	7802		7946	9	10083	10244	10571	2149	4922	2586	8475	9074	10176	10632	

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| Top Hit Descriptor | H MCS0003782 Human adult (N. 2) - 1-3 T | RC2-HT0252-120700 044 L40 Under 1 | Home senions afronces of the control

 | EST68746 Fetal lung II Homo sapiens cDNA 5' end | Human germline T-cell receptor beta chain Doparnine-beta-hydroxylase-like, TRY1, TRY2, TRY3, TGRBV27S1P, TCRBV22S1A2N1T, TCRBV3S1A1T, TCRBV3S1A1N2T, TCRBV6S7P, TCRBV5SA2T, TCRBV13S2, TCRBV6S7P, TCRBV7S3A2T, TCRBV13S2A1T, TCRBV9S2A2PT, TCRBV7S2A1N4T,
 | CABVISCON SAME SA | PM24 INDRA 24/380 AGE CON INDRES IN CORS | Homo sapiens chromosome 21 comment 1527 2000 | Homo sapiens chromosome 21 segment HS21Choes | Homo saviens hypothetical protein (A Eggaden) DNA | Homo saplens eukanotic initiation factor 481 (EllE484)
 | Homo sapiens extendit initiation forth AAI (FIFAAI) gene, partal cds | EST68385 Infant brain Home contact 4A (EIF4A1) gene, partial cds | zw04c08 r1 Scenes NhHMpi. S4 Long control of the co | Vol 1807 81 Shares (etal liver spleam ANET S. U.
 | Home satiens champeome 21 perment Usas Const. | Wm94f10x1 NCI CGAP Ut2 Home conjune a live of a conjune of the original of the | 234412.1 Sogres_NhHMPu_S1 Homo sepiens cDNA clone IMAGE:665278 5' similar to gb:L07807 DYNAMIN-1 (HUMAN);

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 | hu09e08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3168120 3' similar to contains MER18.t3 | PROTEIN MOV-10
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Single Exon Probes Expressed in Brain	Top Hit Descriptor	hx80e02.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:3194090 3' similar to TR:055091	Homo saplens chromosome 21 segment HS21 Ch47	772c08x1 Sogres NSF F8 9W OT PA P ST Home seniors of NA plans NA OF progress of	7772c08.x1 Soares, NSF F8 9W OT PA P S1 Home sariens child claractive of	H. sapiens PADPRP-I gene for NAD/+1 ADP-ribns/dransferase	Homo saplens chromosome 21 segment HS21CnR4	DKFZp761B1710 r1 761 (swenym hamy) Home seriors cDNA class DKFZ-Toxp6220 21	258.1 KDA PROTEIN C210RF6 (KIAA0933)	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 2 PRECI IRSOR	Human transposon-like element mRNA	of47b09.s1 Sogres testis NHT Homo saplens cDNA clone IMARR ਜਾਮਕਾਰਕਰਾ ਨਾ	Homo saplens chromosome 21 segment HS21C049	EST66142 Kidney IX Homo sapiens cDNA 5' and similar to EST containing 1' repeat	2x83h08.r1 Sogres_total_fetus_Nb2HF8_9w Homo sepiens cDNA clone IMAGE:799187 5' similar to contains	52411 Himan refine cDNA rendership of the	MR1-CT0352-240200-105-M8 CT0352 Home control of the CT0352-240200-105-M8 CT0352-240200-105-M8	Homo saplens Xa pseudosi tracion, someni 112	Homo saplens chromosome 21 segment HS21Cn48	H.saplens PADPRP-I gene for NADI+) ADP-ribosultransferase	nc11c02.r1 NCI_CGAP_Pr1 Homo sepiens cDNA clone IMAGE:1007810 similar to contains Alu repetitive element:	Homo saplens shox dene, alternativaly splined products complete and	2d79d03.s1 Soares_fetal_heart_NbHH19tW Homo sepiens cDNA close IMAGE:346853 3' similar to	HOMAN);	Homo septems CCAAT-how-binding transcription factor (CBF2) mRNA	Homo sapiens 959 kb contin balween AMI 1 and CRP4 An chromosome 22.22	Homo sapiens basic transcription factor 2 p44 (btf2p44) gene, partial cds, neuronal apoptosis inhibitory profein (nain) and eurold mode an	Homo segiens nucleolar phospharatein B23 (Albita) — BN/A	Homo saplens nucleolar phosphorotal B23 (NPM) mrNA, complete cds	Control of the prospring in the prospring of the prosprin
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	Expression Signel	1.19	1.7	4.8	4.8	2.43	7.99	7.46	1.1	3.01	0.85	0.65	0.67	0.93	8.4	0.68	0.62	1.78	1.62	22.07	2.41	1.75	0.72	2.01	2.01	1.17	1.59	3.25	3.25	1
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	Probe SEQ ID NO:	7800	10147	10945	10945	कू	1236	1655	2328	3916	92/9	2033	88	96882 8	7351	7423	7717	8612	11233	12428	12495	12834	974	1087	1087	1630	2892	2926	2926	7000

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Top Hit Descriptor	2n35b03.s1 Scares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:414029 3' similar to contains Alu repetitive element,contains element MER22 repetitive element;	Homo saplens chromosome 21 segment HS21C083	Human breakpoint cluster region (BCR) gene, complete cds	CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS)	wd39b05.x1 Soares NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2330481 3' similar to contains MER25.t1 MER25 repetitive element;	Hamo seplens chramosome 21 segment HS21 C083	Homo sapiens chromosome 21 segment HS21C083	Homo saplens GTP binding protein 1 (GTPBP1), mRNA	ye24e05.r1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:118688 5'	MR0-SN0040-050500-002-c07 SN0040 Homo sapiens cDNA	we78h03.x1 Soares Dieckgraefe_colon_NHCD Homo sapiens cDNA clone IMAGE:2347253 3; sImilar to SW:RL29 HUMAN P47914 60S RtBOSOMAL PROTEIN L29; contains element PTR5 repetitive element;	1448b09.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo saplens cDNA clone IMAGE:2144537 3' similar to TR:000372 000372 PUTATIVE P150.:	Homo sapiens MCM4 (MCM4) and DNA-PKcs (PRKDC) genes, partial cds	QV1-BT0631-150200-071-f01 BT0631 Homo sapiens cDNA	EST89564 Small intestine I Homo sapiens cDNA 5' end	Homo sapiens lens major intrinsic protein (MIP) gene, complete cds	Homo saplens TPA inducible protein (LOC51586), mRNA	Homo sapiens TPA inducible protein (LOC51589), mRNA	LYSP100 PROTEIN (LYMPHOID-RESTRICTED HOMOLOG OF SP100)	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	H.saplens DHFR gene, exon 3	EST51247 Gall bladder II Homo saplens cDNA 5' end	II.3-HT0619-110700-209-D12 HT0619 Homo sapiens cDNA	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT	Homo sapiens presentin-1 gene, exons 1 and 2	Homo sapiens presentlin-1 gene, exons 1 and 2	Homo sapiens MADS/MEF2-family transcription factor (MEF2C) mRNA, complete cds
Top Hit Database Source	EST_HUMAN	F	NT	SWISSPROT	EST HUMAN	N N	N.	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	N	TN	TN	SWISSPROT	SWISSPROT	SWISSPROT	TN	EST_HUMAN	EST_HUMAN	SWISSPROT	NT	NT	님
Top Hit Acession No.	AA719297.1	AL163283.2	U07000.1	P26694	Al688474.1			11418127 NT	T93176.1	AW867740.1	AI870071.1	A1452982.1	U63630.2	BE080748.1	AA376832.1	U36308.2	7706225 NT	7706225 NT	Q13342	P08548	P08547	X00856.1	AA345220.1	BF352883.1	P35084	AF029701.2	AF029701.2	L08895.1
Most Similar (Top) Hit BLAST E Value	1.0E-09		1.0E-09	1.0E-09	1.0E-09	1.0E-09	1.0E-09	1:0E-09		9.0E-10	9.0E-10	_			8.0E-10	8.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	
Expression Signal	6.4	99.0	1.89	3.13	0.85	2.92	1.68	2.25	1.35	3.74	4.41	4.78	13.27	0.88	3.17	2.44	9.38	9.38	2.24	3.17	24.23	2.19	4.18	1.08	1.48	1.6	1.6	1.67
ORF SEQ ID NO:		30923	L	31795	34124	L		30717		26707	28256		L	L				26108	L			28491	31836	33078		33696	23697	
Exan SEQ ID NO:	17476	18215	L	l	20983	ı	ı	(24857	1	15606	ì	1_	L	16917	22515	13460	13460	14365	14748	15278	15850	18870	19999	20226	20570	20570	Ш
Probe SEQ ID NO:	4744	5416	6740	6053	8280	10212	11789	12333	12503	1286	2838	6735	142	3337	4177	9865	685	685	1618	2013	2564	3085	6092	7316	7556	7875	7875	10209

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Probe	ű.			Most Similar	15 	igle Exon Pro	Single Exon Probes Expressed in Brain
SEQ IS	<i>o</i>	ORF SEQ ID NO:	Expression Signal	(Top) Hit BLAST E Value	Top Hit Acession No.	l op Hit Database Source	Top Hit Descriptor
893			3.5	6.0E-1	0 AJ400877.1	Ĭ	Homo sapiens ASCL3 gene, CEGP1 gene, C11orf14 gene, C11orf15 gene, C11orf16 gene and C11orf17 gene
2884	lj	28132	1.21	6.0E-10	0 Al424405.1	EST HUMAN	HOZDOTX1 NCI CGAP Prze Homo sepiens cDNA close IMAGE prospos al
4689 889	17423		2.7	6.0E-10	0 AW853719.1	11. i	RC3-CT0254-031099-012-q12 CT0254 Home saplens cDNA
8682	21374	34518		6.0E-10	0 P33730	SWISSPROT	E-SELECTIN PRECURSOR (ENDOTHELIAL LEUKOCYTE ADHESION MOLECULE 1) (ELAM-1) (LEUKOCYTE-ENDOTHELIAL GELL ADHESION MOLECULE 2) (I FCAMA) (CDROS)
8682		34519	1	6.0E-10	P33730	SWISSPROT	E-SELECTIN PRECURSOR (ENDOTHELIAL LEUKOCYTE ADHESION MOLECULE 1) (ELAM-1) (LEUKOCYTE-ENDOTHELIAL CELL ADHESION MOLECULE 2) (F. CAMANA)
9534		35373	0.46	6.0E-10	10 P98073	SWISSPROT	ENTEROPEPTIDASE PRECURSOR /ENTERORINASE)
11950			2.18	6.0E-10	6.0E-10 AW971923.1	EST HUMAN	EST384012 MAGE resequences, MAGI Homo seniens chiNA
745			7.27	5.0E-10	5.0E-10 AL046804.1	EST HUMAN	DKFZp434N219 r1 434 (synonym) htec3) Home sanjens cDNA chas DKEZAA2AN346 El
3468	_ 1	28878	2.5	5.0E-10 Q01033	001033	SWISSPROT	HYPOTHETICAL GENE 48 PROTEIN
483	Į	30269	1	5.0E-10	5.0E-10 AF181897.1	N	Homo sapiens WRN (WRN) gene, complete cds
7222	J		1.51	5.0E-10	59.1	EST_HUMAN	801822184F1 NIH, MGC 75 Home saplens cDNA clone IMAGE 4042413 FI
9438	J	35288	1.89	5.0E-10 P34678		SWISSPROT	HYPOTHETICAL 67.9 KD PROTEIN ZKESS 8 IN CHROMOSOME III
9436	22114	35289	1.89	5.0E-10 F		SWISSPROT	HYPOTHETICAL 67.9 KD PROTEIN ZK688.8 IN CHROMOSOME III
109	12830	-	1.17	4.0E-10	A1221083.1	EST HUMAN	1990909.X1 Soares_placenta_8to9weeks_ZNbHP8to9W Homo sapiens cDNA clone IMAGE:1789049.3' similar to contains LTR8 ho 1 TR8 monthly a clonest
292	13348	25976	0.74	4.0E-10 /	AA515260.1	EST HUMAN	Inf84e01.s1 NCI CGAP Co3 Home septens cDNA clone IMAGE ocagas 2"
1989	1	27446	1.31	4.0E-10	AW594709.1	EST HUMAN	hg58g03.x1 NC_CGAP_GC6 Homo saplens cDNA clone IMAGE:2949844 3' similar to contains Alu repetitive element:
2580	15294	28032	3.73	4.0E-10	AL163303.2	LN	Homo saplens chromosome 21 segment HS21C103
7076	19767	32831	25.71	4.0E-10/A	AF224669.1	LΝ	Homo septens mannosidase, beta A, tycosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (UBE2D3) genes. complete cas
10095	22743	35957	0.49	4.0E-10	AW283243.1	EST HUMAN	UI-H-BIZ-ahl-a-07-0-UI.s1 NCI CGAP Sub4 Homo saniens cDNA close (MA GE: 2727084 2)
10343	22880	36208	0.89	4.0E-10	AI267342.1	EST_HUMAN	aq83h11.x1 Stanley Frontal SN pool 2 Homo sapiens cDNA clone IMAGE:2035653
895	13663	26329	3.55	3.0E-10 N	N36113.1	EST HUMAN	yy32f06.s1 Seares melanocyte 2NbHM Homo sapiens cDNA clons IMAGE:272963 3' similar to contains
1329	14078		4.72	3.0E-10	1.7	NT	Homo saplens extracellular divoporatein lacrifin precursor years complete add
4 88	17234	29864	1.04	3.0E-10	AL163203.2	N.	Homo saplens chromosome 21 segment HS210003
4498	17234	29865	1.04	3.0E-10 A	AL163203.2	LΝ	Homo saplens chromosome 21 segment HS21C003
888	18169	30855	1.24	3.0E-10 N	1		yz11g08.s1 Soares_multiple_sclerosis_ZNbHMSP Homo saplens cDNA clone IMAGE:2827823'
919	18887	31856	2.52	3.0E-10 P			RHOMBOID PROTEIN (VEINLET PROTEIN)
6258	19032	32007	3.43	3.0E-10 B	BE302970.1	EST_HUMAN	ba76d08.y1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:2906319 5'

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Probe SEQ 1D NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7660		33432	1.42	3.0E-10	0 AV743302.1	EST_HUMAN	AV743302 CB Hamo sapiens cDNA clone CBFBGD08 5'
7660				3.0E-10	3.0E-10 AV743302.1	EST_HUMAN	AV743302 CB Hamo septens cDNA clone CBFBGD08 5
8829	21321	34463	1.2	3.0E-10	0 H87208.1	EST HUMAN	ys74b12.s1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:220511 3' similar to contains MER29 repetitive element:
8847	1	34784		3.0E-10	0 AW850731.1	EST HUMAN	IL3-CT0219-160200-064-B06 CT0219 Homo saplens cDNA
8947	ı	34785	1.58	3.0E-10	0 AW850731.1	EST HUMAN	IL3-CT0219-160200-064-B06 CT0219 Hamo sapiens cDNA
9240	i		0.58	3.0E-1	0 AF020503.1	Z	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5
10359			2.37	3.0E-1	0 T65891.1	EST_HUMAN	yc11e12.r1 Stratagene lung (#537.210) Homo saptens cDNA clone IMAGE:80398 5'
10493	ı		1.34	3.0E-1	0 AA769294.1	EST_HUMAN	nz36g03.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1289908 3'
12584	1	31003	2.65	3.0E-10	0 BE179517.1	EST_HUMAN	L3-HT0618-110500-136-E07 HT0618 Homo sapiens cDNA
34		25479		2.0E-10	0 P48988	SWISSPROT	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B)
34	12862	25480		2.0E-10	0 P48988	SWISSPROT	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B)
1890	14677		1 96	2 OF-1	0 1 180017 1	L-Z	Homo saplens basic transcription factor 2 p44 (btt2p44) gene, partial cds, neuronal apoptosis inhibitory protain frain) and sundyal motor narron protain ferm) cames, complete ods.
2985	1		1.04	20E-1	0 BF675047.1	EST HUMAN	602/36640F1 NIH MGC 83 Homo sapiens cDNA clone IMAGE:4273377 5
5714	18507		2.54	2.0E-1	0 028640	SWISSPROT	(HPRG)
						,	Homo sapiens cytochrome P450 polypeptide 43 (CYP3A43) gene, partial cds; cytochrome P450 polypeptide 4 (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A7) genes, complete cds; and cytochrome P450
6156		31900	1.37	2.0E-10	0 AF280107.1	F	polypeptide 5 (CYP3A5) gene, partial cds
7279		33039	6.47	2.0E-10	0 BE7910821	EST_HUMAN	601586208F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3940824 5'
7912	20607	33737	.0.48	2.0E-10	0 P26809	SWISSPROT	POL POLYPROTEIN (CONTAINS: PROTEASE; REVERSE TRANSCRIPTASE; RIBONUCLEASE H]
7912		33738	0.48	2.0E-10	0 P26809	SWISSPROT	POL POLYPROTEIN (CONTAINS: PROTEASE; REVERSE TRANSCRIPTASE; RIBONUCLEASE H]
0000	24874		000	200.40	DE434565 4	ECT LINAM	7078d08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3842303 3' similar to contains L1.t3 L1
11297	1		1.37	2.0E-10	0 A 1862153.1	EST HUMAN	1970712.x1 Sognes total fetus Nb2HF8 9w Homo sepiens CDNA clone IMAGE: 2043695.31
1498	1		1.87	1.0E-10	AW867767.1	EST HUMAN	MR0-SN0038-290300-001-01 SN0038 Homo sepiens cDNA
1602	1	27037	3.18	1.0E-10	0 AV652123.1	EST HUMAN	AV652123 GLC Homo sepiens cDNA clone GLCCXA113'
2586	15300		3.16	1.0E-10	0 AW852001.1	EST_HUMAN	QV0-CT0225-191199-058-e08 CT0225 Homo saplens cDNA
3491		28901	0.89	1.0E-10	0 AW832912.1	EST_HUMAN	QV2-TT0003-161199-013-g10 TT0003 Homo saplens cDNA
3528			0.7	1.0E-10		EST_HUMAN	DKFZp434N1317_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N1317 5
3825	16284		1.03	1.0E-10	AL041685.1	EST_HUMAN	DKFZp434N1317_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N1317 5
3996	16744		6.19	1.0E-10	AF213884.1	NT	Homo saplens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete cds

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Top Hit Descriptor	tm54c09.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2161836 3'	yw48e06.s1 Weizmann Olfactory Epithelium Homo sapiens cDNA clone IMAGE:256298 3'	x/45h11x1 NCI_CGAP_Bm50 Homo sapiens cDNA done IMAGE:2621061 3' similar to contains MER10.t1	MEKTO repetitive element;	ES I 34382 Embryo, 6 week I Homo saplens cDNA 5' end	Homo saplens WEE1 gene for protein kinase and partial ZNF143 gene for zinc finger transcription factor	Homo sapiens SNCA isoform (SNCA) gene, complete cds, alternatively spliced	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE ; ENDONUCLEASE)	Human matrix Gla protein (MGP) gene, complete cds	Human matrix Gla protain (MGP) gene, complete cds	Homo sapiens chromosome X region from filamin (FLN) gene to glucose-6-phosphate dehydrogenese	GO'N) gene, complete cass	MET REVERSE INVIOLENT ASE HOMOLOG	AV727859 HTC Homo saplens cDNA clone HTCASC06 5'	CM0-BT0281-031199-087-a03 BT0281 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C083	Hano sapiens chranosome 21 segment HS21 C083	ALDEHYDE OXIDASE	Homo sapiens chromosome 21 segment HS21C013	Homo saplens protocadherin beta 3 (PCDHB3), mRNA	zu01b12.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:730559 5'	601507531F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909295 5'	Hamo sapiens chromosome 21 segment HS21C047	HUMSUPY069 Human brain cDNA Homo sapiens cDNA clone 069	PRE-MRNA SPLICING FACTOR RNA HELICASE PRP2	zv69f10.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:757963 6' similar to TR:G1055250 G1055250 PHEROMONE RECEPTOR VN4.:	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3	(UBE2D3) genes, complete cds	RC1-HT0256-210100-013-f08 HT0259 Homo sapiens cDNA	#82g12.x1 NCI_CGAP_Bm23 Homo sepiens cDNA clone IMAGE:2105830 3' similar to WP:ZK353.1 CE00385;
Top Hit Database Source	EST HUMAN to	EST HUMAN		Т	EST HOMAN	- TN	±N	SWISSPROT B	\ L	Į.		T	7				Т	SWISSPROT A	ΤN			T_HUMAN	노	EST_HUMAN F	Г	EST HUMAN			EST_HUMAN F	EST_HUMAN O
Top Hit Acession No.	AH78617.1	N23712.1		A 00006158.1	AA330642.1	AJ277546.2	AF163864.1	P11369	M55270.1	M55270.1			1		BE063509.1	AL163283.2	AL163283.2	P48034	AL163213.2	11416799 NT	AA436042.1	BE885900.1	AL163247.2	D44668.1	P20095	AA442630.1			BE149425.1	A1609753.1
Most Similar (Top) Hit BLAST E Value	8.0E-11				7.0E-11	7.0E-11	7.0E-11	7.0E-11	6.0E-11	_		_			6.0E-11			5.0E-11	5.0E-11	5.0E-11	4.0E-11		4.0E-11	4.0E-11		4 0E-11	_	_	4.0E-11	4.0E-11
Expression Signal	. 0.7	4.88		0.65	1.75	1.03	2.05	1.17	7.01	7.01		0.67	3.60	7.81	0.52	1.49	1.9	1.36	1.63	14.05	1.94	7.14	1.16	0.85	3.2	0.80		4.5	1.79	0.9
ORF SEQ ID NO:	29334	ļ		1	26862		34224		25837	25838		32398	33308	34095	35059	25451	25451	29571	32187	33194		28238	28385	29939	32153	32686				35402
SEQ ID	16695	16768	ı	_ {	14177	_	21089	77.722	13188	13188						12838	12838	16944	19191	20107	14127	15498	15735	17311	19153	19641	l	_	21962	22215
Probe SEQ ID NO:	3945	4022	į	6574	1430	3852	8396	10129	403	2	3	2220	3	8262	9213	11	3359	4203	6423	7430	1380	2783	2969	4578	6384	8903		7274	. 9295	9562

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ost Similar (Top) Hit Acession Database Source Source	2.0E-11 AA035369.1 EST_HUMAN zk27g02.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:471794 3'	2.0E-11 AA035369.1 EST_HUMAN ZKZ7g02.s1 Soares_pregnant_uterus_NbHPU Homo saplens cDNA clone IMAGE:471794 3'	2.0E-11 AA281956.1 EST_HUMAN Zs18b04.r1 NCL_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:685519 5'	AA704195.1 EST_HUMAN	2.0E-11]AW842143.1 EST_HUMAN RCo-CN0027-210100-011-c01 CN0027 Homo sapiens cDNA	2.0E-11]BF377659.1 [EST_HUMAN CM2-TN0140-070900-372-g01 TN0140 Homo septems cDNA	D25217.2	2.0E-11 P08547 SWISSPROT LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	2.0E-11 11417966 NT Homo sapiens SEC14 (\$. cerevislee)-like 2 (SEC14L2), mRNA	AJ131016.1 NT	AL163279.2 NT	1.0E-11/AF119914.1 INT Homo sapiens PRO3078 mRNA, complete cds		1.0E-11/AF000573.1 NT Homo saplens homogenitisate 1.2-dloxygenase gene, complete cds	1.0E-11 BE004315.1 EST_HUMAN CM0-BN0105-170300-292-d12 BN0105 Homo septens cDNA	1.0E-11/AL163247.2 NT Homo saplens chromosome 21 segment HS21C047		EST_HUMAN	4885546 NT	1.0E-11 R13174.1 EST_HUMAN yr73d08.r1 Soares Infant brain 1NIB Homo saptens cDNA clone IMAGE:28166 5'		BF365119.1 EST_HUMAN	BF680078.1 EST_HUMAN	9.0E-12/AL163300,2 NT Home saplens chromosome 21 segment HS21C100	9.0E-12 AL 163300.2 NT Homo sapiens chromosome 21 segment HS21C100	8.0E-12 BE074720.1 EST_HUMAN IL6-BT0678-130300-038-G12 BT0578 Homo sapiens cDNA	AJ271736.1 NT	Q05904 SWISSPROT	AA704735.1 EST_HUMAN	6.0E-12]AV730554.1 EST HUMAN AV730554 HTF Homo sapiens cDNA clone HTFAWF06 5	nz88f11.s1 NCL_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1302573 3' similar to contains Alu 6.0E-12/AA732516.1 EST HUMAN repetitive element	
Most Similar (Top) Hit BLAST E Value			2.0E-1				2.0E-1		2.0E-1				1.0E-1	1.0E-1				_		1.0E-1	1.0E-1	1.0E-1	1.0E-1	9.0E-1	1-30.6	8.0E-1	8.0E-1	7.0E-1	1-30.7	6.0E-1	6.0E-1;	l o
Expression Signal	1.48	1.48	1.57	1.54	3.54	1.87	2.67	3.14	3.37	1.34	3.35	2.36	1.13	291	1.2	16,93		0.63	3.15	5.44	1.89	1.89	1.62	1.07	1.07	0.93	3.91	1.16	9.59	0.71	8.52	1 0
ORF SEQ ID NO:			37035			31123				26078	26611		27494	27582				31456	33926	34317	34782	34783	37212	35542	35543			29982	37316		2,9678	72000
Exon SEQ ID NO:	Ł		23760	25332	24567	24586	24748	L	25035	13437		14232	14765	14853	16246	L	<u>L</u>		20795	21172	21637			22348		21918	24617	17348	24013	16291	17053	
Probe SEQ ID NO:	11056	11058	11090	12017	12048	12073	12332	12479	12781	661	1195	1485	2030	2122	3490	5249		5741	8101	8480	8946	8946	11257	9697	2696	9237	12125	4613	11322	3535	4314	100

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z01912.s1 Scares_fetal_heart_NbHH19W Homo sapiens cDNA olone IMAGE:375718 3' similar to contains od10g11.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1387588 similar to contains MER29.t2 b28h05.x1 NCI_CGAP_Lu24 Homo seplens cDNA clone IMAGE:2270745 3' similar to TR:Q13539 Q13539 nad21b03.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE;3366077 3' similar to contains MER7.b2 DKFZp434J0428_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434J0426 6*
Homo sapiens Xq pseudoautosomal region; segment 1/2
OLFACTORY RECEPTOR 1D2 (OLFACTORY RECEPTOR-LIKE PROTEIN HGMIP07E) (OLFACTORY Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, Bos taurus Mitch2 mRNA for mitochandrial carrier homolog 2, complete cds. Homo septiens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 3/3. Homo septiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosamal protein (L44L) and FTP3 (FTP3) genes, complete cds. 374g11:s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:480g78 3* 374g11:s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:460g78 3* DKFZp434B1815_s1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434B1815 3 DKFZp434B1615_s1 434 (synonym: https3) Homo saplens cDNA clane DKFZp434B1616 3 EST04462 Fetal brain, Stratagene (cat#836208) Homo sapiens cDNA clone HFBDV33 1242b05.y1 NCI_CGAP_Bm52 Homo saplens dDNA clone IMAGE:2291217 5 Rattus norvegicus Deleted in coloorectal cancer (rat homolog) (D ∞), mRNA Morone sexetilis myosin heavy chain FM3A (FM3A) mRNA, complete cds Top Hit Descriptor EST386850 MAGE resequences, MAGN Homo saplens cDNA RC1-0T0086-220300-011-b07 OT0086 Homo saplens cDNA Homo sapiens Xq pseudoautosomal region; segment 2/2 Homo sapiens chromosome 21 segment HS21C078 Homo saplens chromosome 21 segment HS21C078 Homo sapiens chromosome 21 segment HS21C103 Homo sapiens chromosome 21 segment HS21C102 Single Exon Probes Expressed In Brain RECEPTOR 17-4) (OR17-4) MARINER TRANSPOSASE. L1.t3 L1 repetitive element; MER29 repetitive element MER7 repetitive element partial cds EST_HUMAN EST HUMAN EST HUMAN Top Hiit Database Source EST_HUMAN ST_HUMAN EST_HUMAN EST_HUMAN SWISSPROT EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN 눋 ż 뉟 뉟 op Hit Acession 6.0E-12 BE047779.1 5.0E-12 BE047779.1 5.0E-12 AJ271736.1 5.0E-12 AL163278.2 5.0E-12 AL079581.1 5.0E-12 AL079581.1 5.0E-12 A.1271735.1 5.0E-12 P34982 5.0E-12 AL163303.2 6.0E-12 AL163302.2 5.0E-12 AW974760.1 5.0E-12 AL040739.1 5.0E-12 AL163278.2 AA700326.1 ġ 8.0E-12 AA847898.1 12 ALD40739.1 5.0E-12|AA033745.1 4.0E-12 BF445140.1 AA700326.1 4.0E-12 AF109907.1 4.0E-12 AB042815.1 4.0E-12 AI689984.1 4.0E-12 AJ229043.1 4.0E-12 U78027.1 6.0E-12 (Top) Hit BLAST E Value Vost Simila 4.0E-12 1.04 5.03 6.41 6.41 11.33 0.94 1.33 4.45 4 Expression 28 28 0.96 3.2 8,0 4.03 0.72 2.76 4.2 Signal ORF SEQ ID NO: 34723 29104 28801 31672 32439 33959 32167 32439 34847 35161 36120 36344 25686 25686 29940 34418 36954 SEQ ID 16466 18715 21585 16144 18715 19424 19168 20822 19424 21584 21696 22910 13047 17312 20835 20190 21279 13047 24774 23691 SEQ ID 9374 3713 **88** 1020 3385 8128 5931 633 6933 8566 8883 9006 9323 10262 10468 ġ 2382 7519 4577 12375 11019 8587 8141

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					***	21	Cirgo Evol Flores Explassed III Digiti
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
602	13380	26011	4.27	3.0E-12	3.0E-12 AW341683.1	EST_HUMAN	hd13d01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2909377 3' similar to TR:014517 O14517 SMRP.;
602	13380	26012	4.27	3.0E-12	-12 AW341683.1	EST HUMAN	Ind13d01.x1 Sogres_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2909377.3' similar to TR:O14517 O14517 SMRP.
5084		30421	0.81	3.0E-12	-12 AL163268.2	LN	Homo sapiens chromosome 21 segment HS210068
5365			1.52	3.0E-12	-12 AF111168.2	NT	Homo saplens serine patmitoy/ transferase, subunit II gene, complete cds; and unknown genes
7854	- 1		0.83	3.0E-12	-12 AW854328.1	EST_HUMAN	RC3-CT0255-031099-011-h02 CT0255 Homo sapiens cDNA
8273	J	34109	0.51	3.0E-12 035453	035453	SWISSPROT	SERINE PROTEASE HEPSIN
9004		34844	0.52	3.0E-12	12 035453	SWISSPROT	SERINE PROTEASE HEPSIN
10551	23247	36483	3.03	3.0E-12	.12 U37672.1	N.	Human prostate specific antigen gene, 5' flanking region
10551		36484	3.03	3.0E-12	3.0E-12 U37672.1	LN L	Human prostate specific antigen gene, 5' flanking region
1649		27084	1.39	2.0E-12	12 AW802131.1	EST HUMAN	ILE-UM0071-120400-065-805 UM0071 Homo ganiens cONA
4094	16836	29462	0.91	2.0E-12	12 J01884.1	F	Rat U3A small nuclear RNA
4094		29463	0.91	2.0E-12	2.0E-12 J01884.1	LN	Rat U3A small nuclear RNA
4387	17124		2.03	2.0E-12	2.0E-12 BE063509.1	EST HUMAN	CM0-BT0281-031199-087-e03 BT0281 Homo sapiens cDNA
4840		30192	1.18	2.0E-12	12 070306	SWISSPROT	TBX15 PROTEIN (T-BOX PROTEIN 15)
4840	17570	30193	1.18	2.0E-12	12 070306	SWISSPROT	TBX15 PROTEIN (T-BOX PROTEIN 15)
-							RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE
Sara	- 1	30491	0.77	2.0E-12		SWISSPROT	ENDONUCLEASE
6385	J		2.8	2.0E-12	57.1	EST_HUMAN	EST383946 MAGE resequences, MAGL Homo sapiens cDNA
7075	-)	32830	3.74	2.0E-12	12 T08169.1	EST_HUMAN	EST06060 Infant Brain, Bento Soares Homo semens cDNA clone HIBBA13 5' end
7244	J	33005	1.02	2.0E-12	12 BE173035.1	EST_HUMAN	MR0-HT0559-200400-015-e08 HT0559 Homo sapiens cDNA
7558		33331	2.2	2.0E-12	11422229 NT	LΝ	Homo sapiens Ac-like transposable element (ALTE), mRNA
9208	ı		1.84	2.0E-12/		LN	Homo saplens putative BPES syndrome breakpoint region protein gene, complete cds
9885	22535		11.12	2.0E-12	12 BE165980.1	EST_HUMAN	MR3-HT0487-150200-113-g01 HT0487 Hamo saplens cDNA
10412	23058	38278	78.0	7000	4 100 4 100 4		990702.x1 Sogres_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1931835 3' similar to TR:Q13538
12032	L		2 84	200-12/		ES - TOWER	GISSO CREZ. FUNCTION UNKNOWN.
2000	1.	+	10,7	Z.UE-12	ı	Z	none saplens chromosome 21 segment HS21C083
12223	24680		2.5	2.0E-12	11418248 NT	LZ	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA
- 4	- 000	- 0					hh90a09.x1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE: 2970040 3' similar to contains MER18.t1
118	12938	25579	2.21	1.0E-12	12 AW627674.1	EST_HUMAN	MER18 repetitive element;
1980	14718		1.39		2 AI871726.1		wm51f07.x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:2439493 3' similar to contains L1.b3 L1 repetitive element;
3067	15833	28476	1.29	1.0E-12	L	N	Homo sapiens testis-specific Testis Transcript Y 2 (TTY2) mRNA partial cds

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Probe SEO ID	Exon	ORF SEQ	Expression	Most Similar (Top) Hit	Top Hit Aces	Igle Exon Pro	Single Exon Probes Expressed in Brain Top Hit
O V		Ö Ö Ö	Signal	BLAST E Value	o Z	Database Source	Top Hit Descriptor
3067	1		1.29	1.0E-12	12 AF000991.1	Į.	Homo saplens testis-sopporing Teasonal V 2/TTV2/ mobils
3855	- 1	29242	26.43	1.0E-12	12 AU132248.1	EST HUMAN	AU132248 NT2RP3 Homo seniers CDNA close NT2BD2304472 E1
3855	16605		26.43	1.0E-12	1.0E-12 AU132248.1	EST HUMAN	AU132248 NT2RP3 Hams capteris COTA successive Control of Control o
5877			2.25	1.0E-12	12 U82828.1	L	Homo sanjens stavla telenciaciaciaciaciaciaciaciaciaciaciaciaciac
5950	18732		1.93	1.0E-12	12 Q9Y2G7	SWISSPROT	HYPOTHETICAL ZINC FINGER DEOTEIN VIA AGG
6438	19206	32202	5	, O	V C2000CT V	ļ	Mus musculus WNT-2 gene, partial cds; putative ankyrin-related protein and cystic fibrosis trensmembrane
7016		32764	2.07	1.0E-12	12 AF196864.1	E L	conductance regulator (CFTR) genes, section 1 of 2 of the complete cds; and unknown gene Homo sanlens nutritive RDES medicine had been asset to the complete cds; and unknown gene
7050	19741	.32802	11.32	1.0E-12	12 A1248533.1	EST HUMAN	dh86404.x1 Soares_fetal_liver_splean_thFLS_S1 Homo septens cDNA done IMAGE:1849614.3' similar to gb:M19603 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (HUMAN);contains MER10.t1 MER10 repetitive element:
7050		32803	11.32	1.0E-12	2 Al248533 1	TOTAL TOTAL	qh66a04.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1849614.3' similar to gb:M1e503 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (HUMAN);contains MER10.t1 MER10
8606	21298	34442	1.16	1.0E-12	2 AA782323.1	EST HUMAN	ac26d05.s1 Strategene overv (#027217) Home control in the control
11273	23834		1.72	1.0E-12		EST HUMAN	he38f07.x1 NCI_CGAP_CML1 Home septems cDNA clone IMAGE:2921317 3' similar to contains element. I TR3 repetitive element.
11942	24497	37809	4.54	1.0E-12	12 AW962164.1	EST HUMAN	EST374237 MAGE resentences MAGG Homo socione - DNA
12150	24637		1.52	1.0E-12	Г	Т	W33h08.x1 NCI CGAP Co16 Homo septems CDNA plans IMA CE popular of
12284			2.92	1.0E-12		Т	Homo sapiens chromosome 21 segment HS21Cn68
3618	- 1		+	9.0E-13			Homo sapiens Xq pseudoautosomal region; segment 1/2
/788	- 1	29320	0.98	9.0E-13 /	5.1		Homo saplens CST gene for cerebroside sulfotransferase, exon 1, 2, 3, 4, 6
LOCA C	-		2.67	9.0E-13		EST_HUMAN	za26b06.s1 Soares fetal liver spieen 1NFLS Home sablens cDNA clone IMAGE:203861.3
8	13475	26123	7.37	8.0E-13 (TN	Homo sapiens priori protein (PrP) gene, complete cds
8	13475	26124	7.37	8.0E-13	3 U29185.1	NT	Homo saplens prion protein (PrP) gene, complete cds
1830	14569	27281	2.94	8.0E-13 U80017 1		Ŀ	Homo sapiens basic transcription factor 2 p44 (bt/2p44) gene, partial cds, neuronal apoptosis inhibitory
8011	20706	33834	0.76	8.0E-13/	-	T HUMAN	wm31h09 v1 NCT COAD THE Union retired (smn) genes, complete cds
8011	20706	33835	0.76	8.0E-13 /		Т	wm31h09 x1 NCI COAP 114 Home septems of the company
10046	22694		3.08	8.0E-13 U78027.1			Homo septiens Bruton's tyrosine kinase (ETK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) denes, complete cds.

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Top Hit Descriptor	Human germline T-cell receptor beta chain TCRBV13S1, TCRBV6S8A2T, TCRBV5S6A3NZT, TCRBV13S6AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6S9AZT, TCRBV6SAAZT, TCRBV6S4A1, TCRBV6SAA	wm31h09.x1 NCI_CGAP_UM Homo sapiens cDNA clone IMAGE:2437601 3'	wm31h09x1 NCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:2437601 3'	OLFACTORY RECEPTOR-LIKE PROTEIN OLF2	601463285F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3866613 5'	POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE) (UDP-GALNAC:POLYPEPTIDE, N-	ACETYLGALACTOSAMINYLTRANSFERASE) (GALNAC-T1)	Homo sapiens chromosome 21 segment HS21C007	y82/04.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:145759 5'	277a12.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728350 3' similar to contains Alu	repetitive element; contains element MER22 repetitive element;	GAP JUNCTION BETA-1 PROTEIN (CONNEXIN 30) (CX30)	MYOSÍN LIGHT CHAÍN KINASE, SKELETAL MUSCLE (MLCK)	PM2-HT0224-221099-001-e11 HT0224 Homo sapiens cDNA	Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	PM3-HT0520-230200-002-c08 HT0520 Homo sapiens cDNA	Homo saplens mRNA for KIAA1329 protein, partial cds	zw76g12.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:782182 5' similar to TR:G452763 G452763 COR1 MRNA.;	yy33g05.r1 Scares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:273080 5' similar to PIR:A32895	A32995 t complex stertlity protein - mouse;	DKFZp434A0128_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434A0128 6'	7804H11 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7B04H11	qr32d05,x1 NCI_CGAP_Kid5 Homo saplens cDNA clone IMAGE:1899945 3' similar to contains Alu	repetitive element;	z/78g10.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728514 3'	z/78g10.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:7285143'	Homo sapiens X-linked anhidrotito ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	zw68g08.r1 Scares_testis_NHT Homo sapiens cDNA clone IMAGE:781406 51
Top Hit Database Source	E	EST HUMAN	EST_HUMAN	SWISSPROT	EST_HUMAN		SWISSPROT	N	EST_HUMAN		EST_HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN	LN	EST_HUMAN	NT	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	ΤN	EST_HUMAN
Top Hit Acession No.	3 U68060.1	3 AI884398.1	3 AI884398.1	3 Q95155	3 BE778223.1		3 Q10473	3 AL163207.2	3 R78338.1		3 AA435773.1	3 P08983	3 P07313	3 AW378614.1	3 AF003529.1	3 BE169131.1	3 AB037750.1	3 AA431528.1		4.0E-13 N44291.1	3 AL043810.1	3 AA076907.1		4.0E-13 AI289831.1	4.0E-13 AA435819.1	4.0E-13 AA435819.1	3.0E-13 AF003528.1	3.0E-13 AA430310.1
Most Similar (Top) Hit BLAST E Value	8.0E-13	7.0E-13	7.0E-1	7.0E-1	7.0E-1		7.0E-13	6.0E-13	5.0E-13		5.0E-13/	5.0E-13	5.0E-13	4.0E-13	4.0E-13	4.0E-13	4.0E-13	4.0E-13		4.0E-13	4.0E-13	4.0E-13		4.0E-13	4.0E-13	4.0E-13 /	3.0E-13	3.0E-13/
Expression Signal	1.67	0.71	0.71	99.0	3.05		1.37	6.75	0.74		1.54	0.84	2.72	2.23	1.67	5.51	1.05	0.94		1.07	1.07	0.45		4.94	2.09	2.09	4.94	1.62
ORF SEQ ID NO:	37701		33496					27558				32548	36693			31195	32858	33277							37066			
Exan SEQ ID NO:	24370	<u>L</u>			24788		_	14825	16078	Ì			23451		15180		19793	20183	1	20286	21432	22084			23789	23789	12987	13615
Probe SEQ ID NO:	11778	7718	77.18	8133	12404		12617	2094	3316		3392	67777	10767	1860	2462	5499	7105	7512		7620	8740	9402		9919	11120	11120	175	845

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Table 4
Single Exon Probes Expressed in Brain

Тор Hit Descriptor	CELL SURFACE GLYCOPROTEIN 1 PRECURSOR (OUTER LAYER PROTEIN B) (S-LAYER PROTEIN 1)	S.scrofa rps12 mRNA for ribosomal protein S12	Human PFKL gene for liver-type 8-phosphofructokinase (EC 2.7.1.11) expn 2	Homo sapiens N-myristoy/transferase 1 (NMT1), mRNA	Homo saplens N-myristoyftransferase 1 (NMT1), mRNA	Homo saplens mab-21 (C. elegans)-like 1 (MAB21L.1) mRNA	CM0-NN0001-100300-274-e11 NN0001 Homo sepiens cDNA	FGF-1=fibroblast growth factor 1 [human, kidney, Genomic, 342 nt, segment 2 of 2]	Homo sapiens LGMD2B gene	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RINGB, 9, 13 and 14	nw21g02 s1 NCI CGAP GCB0 Homo septens cDNA clone IMAGE:1241138 3' similar to contains THB t3	THR repetitive element;	602038009F1 NCI_CGAP_Bm64 Homo sepiens cDNA clone IMAGE:4185866 57	nn24d01.s1 NCI_CGAP_Gas1 Homo sapiens cDNA done IMAGE:1084801 3' similar to contains Alu	repetitive element:contains element MER24 repetitive element;	nn24d01.s1 NCI_CGAP_Gas1 Homo sepiens cDNA done IMAGE:1084801 3' similar to contains Alu repetitive element contains dement MER24 repetitive element:	MELANOMA-ASSOCIATED ANTIGEN B4 (MAGE-84 ANTIGEN)	Mus musculus osteofesticular protein tyrosine phosphatase mRNA, complete cds	745e10.xt Soares_NSF_F8_9W_OT_PA_P_S1 Homo saptens cDNA clone IMAGE:3524443 3' similar to contains MER29.b2 MER29 repetitive element:	AV715377 DCB Homo sapiens cDNA clone DCBAIE03 5'	Homo sapiens Xq pseudoautosomal region; segment 1/2	a 24c01.s1 Soares_testis_NHT Homo sapiens cDNA clone 1391232 3' similar to contains MER19.r1 MER19	repetitive element;	ej24001.s1 Soares_testis_NHT Homo sapiens cDNA clone 1391232 3' similar to contains MER19.H MER19	DOL PT0000 0000 000 CT0000 Home conject DNA	CT-C10222-000 C10522 Adillo Sapiens CDINA	Homo sapiens mRNA for sodium-glucose cotransporter (SGLT2 gene)	Homo sepiens mRNA for sodium-glucose cofransporter (SGLT2 gene)	Homo sapiens TFF gene cluster for trefoll factor, complete cds	хо54h05.x1 NCl_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2707833 3'
Top Hit Database Source	SWISSPROT	NT (- N				T_HUMAN		Z	<u> </u>		EST HUMAN	EST HUMAN	Г	EST_HUMAN IN	EST HUMAN	t⊤	LN	EST HUMAN	Т	- L	Г	EST HUMAN		Т	NEWDE				EST_HUMAN
Top Hit Acession No.	13 Q06852			10835072 NT	10835072 NT	5031896 NT	3 AW892155.1	S74129.1	3 AJ007973.1	3 X87344 1		3 AA720574.1	3 BF340987.1		3 AA577812.1	3 AA577812.1		3 AF300701.1	3 BF108755 1	l	3 AJ271735.1		4 AA781159.1	A A 704450 4	1	T				4 AW513298.1
Most Similar (Top) Hit BLAST E Value	2.0E-13	2.0E-13	2.0E-13	2.0E-13	2.0E-13	2.0E-13	2.0E-13	l m	1.0E-13	4 0F-43	21-12:1	1.0E-13	1.0E-13		1.0E-13	1.0E-13	1.0E-13	1.0E-13	1 0F-13	1.0E-13	1.0E-13		9.0E-14	77 300	9.0E-14	8.0E-14	9.0E-14	9.0E-14	9.0E-14	9.0E-14
Expression Signal	4.7	0.58	7.15	0.65	0.65	3.87	3.48	1.52	5.64	40,4	3	2.13	1.64		0.78	0.78	0.79	0.53	11.1	2.25	2.12		1.81	90 0	3,00	8.5	1.18	1.18	2.6	3.96
ORF SEQ ID NO:	31772		32675					25732	26307	82786		27478	29917		33626	33627		36063	37345				25781	05700	70/07		28050	28051	28206	28513
Exen SEQ ID NO:	18812	1		19436	L	L.	L		13637	14081	1	14750	1		20202	20505		1	24042	24492	24893		13125	90,404	15120	125	15313	15313	15463	15874
Probe SEQ ID NO:	6032	6113	6717	6954	6954	-10355	12105	285	888	1313		2015	4553		7810	7810	0668	10199	11352	11934	12563		324	300	250	200	5269	2599	2758	3109

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Sizo Exercise Exercise Copp. Sizo Signal Top-Hit Top				6	_	Г	Т	Т	Г	1	Γ-	$\overline{}$	7-	_	_						_			_						
December Property bes Expressed in Brain		ai24001 st Spares herits Nut us	repetitive element;	Human DNA, SINE repetitive element	Saguinus cedipus gene for seminal vesicia secretaria.	hz71c09.x1 NCI_CGAP_Lu24 Home services contact in the contact in t	W72e03.r1 Soares placenta Nb2HP Homo saviens CDNA Committee 11.13213424 3'	H.sapiens DNA for endogenous retroviral like element	2417c10.s1 Stratagene fetal retine 937202 Homo saniens CONA size 1111.55	QV2-BT0258-281099-014-a01 BT0258 Homo sanians CDNA	We92h08.x1 NCI_CGAP_CG3 Homo saplens cDNA clone IMAGE:2326143 3/	MER10 repetitive element:	Homo sepiens chromosome 21 segment HS210005	Home confirm that the	orino septens FrA35 common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens FRA3B common fragile region, diadenosine triphosohate hydrolasa (FLIIT)	SSOCIATED PROTEIN SOCIATED PROTEIN SOCIATED PROTEIN 1 (MULTIDRUG RESISTANCE	bosbos x1 NCI_CGAP_GU1 Home saniens CDNA class 114 ACE TROTEIN)	spetitive element;	INE-1 REVERSE TRANSCRIPTASE HOMOLOG	-ANTIGEN PROTEIN PRECURSOR	omo saplens LGMD2B gene	K67a06.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMA DE: 1022aze #1	7/3c12.s1 Soares_multiple_sclerosis_2NbHMSP Home saplens cONA clone IMAGE:278180 3' elmillor to	saplens DMA, DMB, HLA-21, IPP2 1 MP2 TABY 1 MP3 TABY 1	anes ', Livit', IAFK, DOB, DQB2 and RING8, 9, 13 and 14	recovery INCL CGAP Out Homo seplens cDNA done IMAGE:24353323' similar to contains Alu	R.narveglous mRNA for CPG2 protein	
Dec Exon CAF SEQ Expression CTop) Hit Top Hit Acess No. No. Signal BLASTE No. Value BLASTE S.B. B.CE-14 AA781159.1 B.C. AA7	gle Exon Pro	Top Hit Database Source		EST_HUMAN	L	LN	EST_HUMAN	EST_HUMAN	LN LN	EST_HUMAN	EST_HUMAN	EST HUMAN	ST HUMAN	5						Г	╗	\Box	ISSPROT	T	Т					П
Dec Exam ORF SEQ Expression No. ID SEQ ID ID NO. Signal ID NO. I	riS.	Top Hit Acess No.		AA781159.1	0.14547.1	AJU02153.1	BE468263.1	I	T	1	T												1	T					-	
Exam ORF SEQ Express		Most Similar (Top) Hit BLAST E Value		1	1	9.0E-14	0.0E-14	0.0E-14	8 OF 14	B OF 14	8 OF 14		7.0E-14	(.0E-14)	6.0E-14	A 1			4	7	5 OF 44 BY	4 0F-14 P	4 OF 44 A	4.0E-14 A		-	_			
De Exon NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:		Expression Signal	100	8 8 A	1.68	1 27	287	15.04	3.69	1.72	2.48		4.77	10.07	14.14	. 6	90		5.46	141	5.77	2.18	5.8	0.87		1.05	0.59		7.02	8
20 C C C C C C C C C C C C C C C C C C C					L	L			35180		31056		1	+	25797	35572	35573		28014	30322	31133		27319		_	29630		-	28381	1
				L			Ι.	J				!	1		13158	22373	22373		13382	17716	18245	15560	14608	16488		17000	20563	25414	13697	
		Probe SEQ II NO:	3232	3778	4707	3488	3837	9348	9460	11410	12302	1625	8818		328	9722	9722	į	3	4993	844 ₆	1101	1870	3736	- 5	4269			1	J

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8EQ ID SEQ ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:	ORF SEQ ID NO: ID Signal 0.92 0.92 0.97 0.97 0.97 0.97 0.97 0.97 0.97 0.97	3.0E-14 3.0E-14 3.0E-14 3.0E-14 3.0E-14 3.0E-14 2.0E-1	Top Hit Acession No. No. AW265354.1 ES. 7656864 NTT AL420786.1 ES. AL420786.1 ES. AL420786.1 ES. AL420786.1 ES. AL420786.1 ES. AL420786.1 ES. AL43228.2 NT AL5328.2 NT AL5328.2 NT AL5328.2 NT AL5328.2 NT AL63303.2 NT AL163303.2 NT AL163303.2 NT AL163303.2 NT AL163303.1 ES. AW372868.1 ES. PO8548 SW BE222432.1 ES. U01317.1 NT BED00550.1 ES.	Tep Hit Database Source THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN THUMAN	Top Hit Descriptor repetitive element; contains element MERe repetitive element; the force element; contains element MERe repetitive element; the force element; contains element MERe repetitive element; the force element; contains element MERe repetitive element; the force element; contains element MERe repetitive element; the force element; contains element MERe repetitive element; the force of the force	
	Ш		20E-14	P56163	SWISSPROT	ZING-FINGER PROTEIN NEURO-D4
7407 20084 7407 20084	33167	22.12	2.0E-14 2.0E-14	BE158761.1 BE158761.1	EST_HUMAN	II.2-HT0397-071299-024-D04 HT0397 Homo sapiens cDNA II.2-HT0397-071299-024-D04 HT0397 Homo sapiens cDNA
ll			2.0E-14	AI978795.1 AV741648.1	EST_HUMAN	wr59g10.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2492034 3' similar to contains Alu repetitive element; AV741648 CB Homo sapiens cDNA clone CBFBBF04 5'

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		T	τ	7	_	_	_				_	<u> </u>	_	-	_							_								
Single Exon Probes Expressed in Brain	Top Hit Descriptor	UI-H-BI1-edw-e-10-0-UI.s1 NCI CGAP Sub3 Homo sepiens cDNA clone IMAGE:2718234.31	xc36f02.x1 NCI_CGAP_Co20 Hamo sapiens cDNA clone IMAGE:2586363 3' similar to contains MER1.t3 MFR1 renalitive element	Homo sablens butative G6 protein (ARR) dene commiste ede	Hamo saplens chromosome 21 segment HS21C046	Homo saplens chromosome 21 segment HS21C068	Homo sapiens chromosome 21 segment HS21C088	Homo saplens chromosome X region from filamin (FLN) gene to glucose-6-phosphate dehydrogenase	Homo sablens chromasome 21 sagment HS21C103	Homo suplens ribosomal protein L23A (RPL23A) gene, complete cds	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PEHRP.II)	RC2-CT0432-310700-013-a09 1 CT0432 Homo sapiens cDNA	RC2-CT0432-310700-013-a09 1 CT0432 Home saplens aDNA	ae39c12.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:971350.3	xq39h10x1 NCI_CGAP_Lu28 Homo saplens cDNA clone IMAGE:27530593'	Bos taurus xenobiotic/medkim-chain fatty acid:CoA ligase form XL-III mRNA, nuclear mRNA encoding	Hong saplens mominin (manes) like 4 (BDON) 11 DNA	Homo sabiens prominin (mouse)-like 1 (PROMI 1) mRNA	RC2-CT0432-310700-013-g09 1 CT0432 Homo saplens cDNA	RC2-CT0432-310700-013-a09_1 CT0432 Homo saplens cDNA	Homo saplens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA	Homo saplens transcription factor IGHIM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T64 protein, IM0 protein, Additional factor of protein, IM1 protein, JM5 protein, T64 protein, IM1 protein, JM2 protein, JM5 protein, T64 protein, JM5 protein,	own by count, art united subdoor region destruction up to Entry domain protein 6, and synaptophysin genes, complete cds; and L-type calcium channel as	GAG POLYPROTEIN CONTAINS: CORE PROTEINS P15, P12, P30, P101	601677750F1 NIH MGC 21 Homo saplens cDNA clone IMAGE: 3960168 6	Homo saplens chromosome 21 segment HS21C047	601148632F1 NIH MGC 19 Homo sablens cDNA clone IMAGE:3164023 5	801458531F1 NIH_MGC_68 Hamo saplens cDNA clone IMAGE:3862086 5	xn77d02.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2700483 3' similar to contains THR.t2 THR repetitive element;	Homo sapiens Xq pseudoautosomal region; segment 2/2
gie Exon Pro	Top Hit Database Source	EST_HUMAN	FST HUMAN	LN	N.	LN	FN	F	LN	LN	SWISSPROT	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	5	F	Į	EST HUMAN	EST_HUMAN	LZ L		L	ISSPROT	Г	Г	EST HUMAN		EST_HUMAN	Ŋ
uio	Top Hit Acesslon No.	-14 AW139800.1	-14 AW083969 1	-14 AF008191.1			-14 AL163268.2	-14 44140 1	22	Γ		14 BF335227.1	14 BF33527.1		14 AW275852.1	14 AF128145 1	17150	11437150 NT	14 BF335227.1	14 BF335227.1	7427522 NT		15 AF186779.1	15 P21416	15 BE903559.1	15 AL163247.2	15 BE261482.1	5 BF035327.1 E	_	
	Most Similar (Top) Hit BLAST E Value	2.0E-14	2.0E-14	2.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.0E-14		1.0E-14	1.0E-14	1.0E-14	9.0E-15		9.0E-15/	9.0E-15	9.0E-15	9.0E-15/	8.0E-15		7.0E-15	6.0E-15
	Expression Signal	4.88	1.29	2.29	1.88	6.41	6.41	12.44	4.55	3.56	1.79	5.42	6.42	1.67	1.91	2.42	11.5	11.5	3.05	3.05	2.08		1.43	4.51	1.08	2.36	0.91	1.14	3.07	8.64
	ORF SEQ ID NO:	36612	37507				26806	27452	27643	27866	28363	28576	28577	29255	29802	31432	32351	32352	28576	28577	27002			33152	33740			32836		26403
	SEQ ID NO:	23370	24190	l			14132	14730					1	ļ l	17176	18511	25095	25095		15928	14317		14899	20074	20610	24991	13253	19771	22981	13738
	Probe SEQ ID NO:	10679	11591	12536	1045	1385	1385	1894	2182	2409	2945	3165	3165	3866	4440	6719	6576	8276	11818	11818	1570		2170	7396	7915	12718	2814	7081	10334	973

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		ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5829	18618		1.02	6.0E-15	5 X73462.1	LN	O.aries mRNA for hair keratin cysteine-rich protein
5829	18618		1.02		X73462.1	N	O.aries mRNA for hair keratin cysteine-rich protein
401	13186	25834	6.63	5.0E-15	5.0E-15 AL163208.2	IN	Homo sapiens chromosome 21 segment HS21C008
2764	15469	28212	1.38	5.0E-15	U91328.1	LN	Human hereditary haemoohromatosis region, histone 2A-like protein gene, hereditary haemoohromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
8 8	16217		1.03	5.0E-15	AW296817.1	EST HUMAN	UI-H-BW0-ajb-g-10-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2731219 3'
10574	23269		2.4	5.0E-15	AV730056.1	EST_HUMAN	AV730056 HTF Hamo saplens cDNA clone HTFAVE08 5'
418	12829	25442	2.85	4.0E-15	AL163303.2	Ĭ	Homo sapiens chromosome 21 segment HS21C103
6567	19332	32339	0.76		AB007970.1	NT	Homo saplens mRNA, chromosome 1 specific transcript KIAA0501
10994	20392	33505		4.0E-15	AJ130894.1	NT	Homo saplens mRNA for transcription factor
10994	20392	33506	3.08		AJ130894.1	IN	Homo saplens mRNA for transcription factor
4192	16933		2.08	3.0E-15	NR9452 1	FST HIMAN	LY1142F Human fetal heart, Lambda ZAP Express Homo saplens cDNA clone LY1142 5' similar to ANF(CARDIODII ATIN)
4872	17599		0.79		P92485	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5
6716	19631		1.33	3.0E-15	Q64625	SWISSPROT	GLUTATHIONE PEROXIDASE RY2D1 PRECURSOR (ODORANT-METABOLIZING PROTEIN RY2D1)
7179	19865	32937	2.9	3.0E-15	M27685.1	LN	Mus musculus ultra high sulfur keratin gene, complete cds
7179	19865	32938	2.9	3.0E-15	M27685.1	LN	Mus musculus ultra high sulfur keratin gene, complete cds
9825	22478		2.51	3.0E-15	AA807128.1	EST HUMAN	oc38e07.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1351784 3' similar to contains MER19.t1 MER19 repositive element
							Homo saniens DNA DI EC1 to ORCIT 4 nene region santon 4/2 (DI EC4 ODCIT) 3 ODCIT 4 minor
10694	23385	36625	2.47		AB026898.1	NT	omplete cds)
12310	25315		1.81	3.0E-15	AJ271735.1	LN	Homo sapiens Xq pseudoautosomal region; segment 1/2
12814	25056		1.35		AW877214.1	EST_HUMAN	CM4-PT0034-180200-506-e01 PT0034 Hamo sapiens cDNA
243	13052	25692	3.6	2.0E-15	AF223391.1	ΤN	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
359	13157	25798	3.90	2.0E-15	AF223391.1	Ļ	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
328	13157	25799	3.99	2.0E-15	AF223391.1	LV.	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, atternatively spliced
3500	16256	28910	0.71	2.0E-15	AF223391.1	TN	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
3500	16256	28911	0.71	2.0E-15	AF223391.1	TN	Homo sapiens calclum channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced

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Single Exon Probes Exonessed in

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oligia Lyoli Floras Lybiassau III Bi alli	Top Hit Descriptor	Homo sapiens hypothetical protein FLJ10024 (FLJ10024), mRNA	AV730030 HTF Homo saplens cDNA clone HTFAWA03 6	FOLLISTATIN-RELATED PROTEIN PRECURSOR	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	Homo saplens Grb2-associated binder 2 (KIAA0571), mRNA	df45c01.y1 Morton Fetal Cochlea Homo sapiens cDNA clone IMAGE:2486376 6'	d/45c01.y1 Morton Fetal Cochlea Homo sapiens cDNA clone IMAGE:2488376 5'	DKFZp434P037_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434P037 5	Hamo saplens TSX (TSX) pseudogene, exon 5	ZONADHESIN PRECURSOR	ENVELOPE GLYCOPROTEIN GP340 (MEMBRANE ANTIGEN) (MA) [CONTAINS: GLYCOPROTEIN GP220]	EST06060 Infant Brain, Bento Soares Homo saplens cDNA clone HIBBA13 6' end	Human BXP20 gene	7B10F02 Chromosome 7 Fetal Brain cDNA Library Homo saplens cDNA clone 7B10F02	Homo saplens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	am98h05.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:1884185 3' similar to contains	IHK.bZ IHR repetitive element;	602246538F1 NIH_MGC_62 Homo sapiens cDNA clane IMAGE:4332032 5'	Homo sepiens ADP/ATP carrier protein (ANT-2) gene, complete cds	Homo saplens chromosome 21 segment HS21C079	af06d04.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1030855 3'	Human SSAV-related endogenous retroviral LTR-like element	H. sapiens DNA for endogenous retroviral like element	995603.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1839197 3' similar to contains MER29.t3 MER29 repetitive element	RC3-BT0048-131199-003-H12 BT0046 Homo sapiens cDNA	HISTIDINE-RICH PROTEÍN KE4	ij16e11.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2141708 3' similar to contains element	MER33 repetitive element;	nz47f06.x5 NCI_CGAP_Pr12 Homo septens cDNA clone IMAGE:1280947 similar to TR:054849 054849 HYPOTHETICAL 42.9 KD PROTEIN [2] TR:008965 :contains MFR7 tr MFR7 reportitive alement	7f82h09.x1 NCI_CGAP_Pr2B Homo saplens cDNA clone IMAGE:3303521 3'
שום ביים	Top Hit Database Source		EST_HUMAN A	SWISSPROT F			EST_HUMAN df	П	EST HUMAN D	Г	SWISSPROT Z	EWISSPROT G		Γ	EST_HUMAN 7E	Ĭ		HOMAN	EST_HUMAN 60	Ĭ.	Ĭ	EST_HUMAN af	NT	NT H.	EST HUMAN ME	Т	Т		EST_HUMAN ME	nz EST HUMAN HY	П
5	Top Hit Acession No.	11423191 NT	B AV730030.1	6 ଦୁହେଥେ	6 P08548	6912459 NT	6 AW022862.1.	6 AW022862.1	6 AL046445.1	6 AF135446.1	6 0 28983	6 P03200	_	6 U03887.1	6 AA077225.1	6 AF003529.1			8 BF690617.1	6 L78810.1		1.1	2.0E-16 J03061.1		20E-16 AI208733.1		2.0E-16 Q31125		AI470723.1	AI732837.1	
	Most Similar (Top) Hit BLAST E Value	4.0E-16	4.0E-18	4.0E-16	4.0E-16	4.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-16	3.0E-18	3.0E-16	07 20 0	3.UE-10	3.0E-16	3.0E-16	2.0E-18	2.0E-16	2.0E-16	2.0E-16	2.0E-16	2.0E-16	2.0E-16		2.0E-16/	2.0E-16/	2.0E-16
	Expression Signal	1.04	1.51	1.44	2.04	2.51	2.03	2.03	1.5	1.5	1.38	3.76	19.63	0.95	0.99	1.79		4.20	0.89	29.9	1.2	0.91	1.06	1.16	96.0	0.79	0.99		0.75	2.14	0.57
	ORF SEQ ID NO:		37133			31087	25589	25590			26867	28388	29304			31230		24201		35883				29526	29807	30439	32419		33389	33689	33883
	Exon SEQ ID NO:						12945	12945	13239	13248	14182	15741	16663	16689	18004	18327	0.70	21540	22441	22667	13717	15106	15403	16897	17183	17822	19404		20281	20562	20752
	Probe SEQ ID NO:	9195	11182	11851	12014	12109	130	130	453	463	1435	2975	3913	3939	5196	6259	0	8	9790	10019	951	2385	2694	4157	4447	5104	6842		7615	7867	8028

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Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
8028	20752	33884	0.57	2.0E-16	6 BE858026.1	EST_HUMAN	7782h09.x1 NCI_CGAP_Pr28 Hamo sapiens cDNA done IMAGE:3303521 3'
8425		34256	0.81	2.0E-16	6 AW877214.1	EST_HUMAN	CM4-PT0034-180200-506-a01 PT0034 Homo sapiens cDNA
8425	ı	34257	0.81	2.0E-16	6 AW877214.1	EST_HUMAN	CM4-PT0034-180200-506-a01 PT0034 Homo sapiens cDNA
8	12992	25630	1.84	1.0E-16	1.0E-16 AF200719.1	LN	Homo saplens pituitary tumor transforming gene protein (PTTG) gene, complete cds
Ę	10100		20.00	1 05 18	6 4 4 6 28 5 0 2 4	EST HIMAN	af39g11.s1 Soares_total_fetus_Nb2HF8_9w Homo saplens cDNA clone IMAGE:1034084 3' similar to contains OFR to OFR renefitive element
1083	14500	27414	23.00	1.0E-16	1.3C-16 RF327942 1	EST HUMAN	OVO-BN0148-070700-283-a10 BN0148 Homo sapiens cDNA
5635	18430	31343	0.75	1.0E-18	1.0E-16 AF163864.1	F	Homo saplens SNCA isoform (SNCA) gene, complete cds, alternatively spliced
6341	19111		27.85	1.0E-16	6 U45983.1	F	Homo sapiens CCR8 chemokine receptor (CMKBR8) gene, complete cds
1	8	9000	0	0.1	000	TOGGOOM	MITOGENACTIVATED PROTEIN KINASE KINASE KINASE 10 (MIXED LINEAGE KINASE 2) (PROTEIN KINASE MST)
27.5	19240	32240	0.08 14 F	100,	0 002778	PA POST NO	Homo contane CORR chemokine receptor (CMKRRR) napa complete ode
3 8	19111	95048	70,		1.0E-10 043905.1	ENT LIMAN	COV2. PTOM 2.2.40400.124.e05. PTOM 2. Home sanlens cDNA
3 3	3017	01000	20.2	١	AW6/3031.1	NONDE FOR	CASE INVITED SOCIETY AS AND MINISTER CONTROL
37.22	16475	27112	2:11	9.0E-17	9.0E-17 AW 900048.1	ESI HOMAN	CAN I-WIN TOUS-ZAUGGO-105-EQ I WIN TOUS FIGHTS CELIAN CONTRACTOR AND CONTRACTOR A
7000	8		Č	74	7 N300084 4	. NAME: 12	tig22c11.X1 NCI_CGAP_CL11 Homo sapiens cUNA ctone IMAGE:2109524 3 similar to contains MEK28.tz MFR28 renotifiue element
5700	13230		7.7	8.0E-17	Aloszao4. I	NAMOR ISS	MILLY CO I OPOUR O CONTROL I
8007	20702		4.75	9.0E-17	7 AW150257.1	EST_HUMAN	xg49g12.x1 NC_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2630950 3' similar to contains OFR.12 OFR repetitive element ;
10124	22772		2.47	9.0E-17	7 AF200719.1	LN.	Homo sapiens pitultary tumor transforming gene protein (PTTG) gene, complete cds
266	13757		1.77	8.0E-17	7 AW880701.1	EST_HUMAN	QV0-OT0032-080300-155-d01 OT0032 Homo sapiens cDNA
3872	16822		0.87	8.0E-17	7 AL163280.2	۲	Homo sapiens chromosome 21 segment HS21C080
5498	25069	31193	3.7	8.0E-17	7 BE172081.1	EST_HUMAN	MR0-HT0559-060300-003-e04 HT0559 Homo sapiens cDNA
7175	19861		<u>4</u> .	8.0E-17	7 AV730759.1	EST_HUMAN	AV730759 HTF Hamo sapiens cDNA clone HTFAQB07 5'
14	14188		3.44	7.0E-17	6753097 NT	TN	Mus musculus apolipoprotein B editing complex 2 (Apobec2), mRNA
5240	18046		3.3	7.0E-17	7 AF216650.1	L	Homo sapiens putative MTAP (MTAP) mRNA, partial cds, alternatively spliced
					į		Mus musculus WNT-2 gene, partial cds; putative ankyrin-related protein and cystic fibrosis transmembrane
6588		32365	8.05	7.0E-17	7 A F 2 2 9 8 4 3 . 1	LN	conductance regulator (CFTR) genes, section 1 of 2 of the complete cds, and unknown gene
198	13011	25653	8	6.0E-17	7 AW983890.1	EST_HUMAN	RC1-HN0003-220300-021-b04 HN0003 Homo saplens cDNA
							hi81d04.x1 Scares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:29786953' similar to contains L1.t2
6221		31971	1.64		6.0E-17 AW662772.1	EST_HUMAN	L1 repetitive element ;
10190	1	36053	0.48	6.0E-1	P20138	SWISSPROT	MYELOID CELL SURFACE ANTIGEN CD33 PRECURSOR (GP67)
412		25438		5.0E-1	7 T64110.1	EST_HUMAN	yc05h08.r1 Strategene lung (#937210) Homo sapiens cDNA clone IMAGE:79839 51
7486	20158	33250	2.09	5.0E-17	7 T81043.1	EST_HUMAN	yd26b04,r1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:109327 5

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Top Hit Descriptor	EST13504 Testis tumor Homo sapiens cDNA 5' end similar to similar to glycogenin	600944690F1 NIH_MGC_17 Hamo sapiens cDNA clane IMAGE:2960615 5'	Homo saplens chromosome 21 segment HS21C047	Homo sapiens chromosome 21 segment HS21C047	Human CYP19 gene for arometase cytochrome P-450, promoter region (containing two cis-acting transcriptional regulatory elements)	BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (BMP-1)	BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (BMP-1)	we94b04.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2348719 3'	we94b04.x1 Scares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2348719 3'	MULTIDRUG RESISTANCE PROTEIN 1 (P-GLYCOPROTEIN 1)	Homo sapiens Xq pseudoautosomal region; segment 2/2	Hamo saplens chromosame 21 segment HS21 C007	COLLAGEN ALPHA 1(III) CHAIN PRECURSOR	Homo sapiens thrombospondin 2 (THBS2) gene, promoter region and exons 1A and 1B	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3	(UBE2D3) genes, complete cds	y30e07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:128388 5'	he38e05.x1 NCI_CGAP_CML1 Homo septens cDNA clone IMAGE:2921312.3' similar to contains Atu repetitive element;contains LTR8.11 LTR8 repetitive element;	qe85b05.x1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:17438253'	qe65b05x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:17438253'	URIDINE PHOSPHÖRYLASE (UDRPASE)	QV0-BT0263-101299-072-d07 BT0263 Homo saplens cDNA	QV3-BN0048-220300-129-c10 BN0046 Hamo saplens cDNA	MYOSIN LIGHT CHAIN KINASE, SMOOTH MUSCLE (MLCK) [CONTAINS: TELOKIN]	248f06.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:795489 3' simitar to TR:G1263081 G1263081 MARINER TRANSPOSASE	2018g12.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:609862 3'	(188403.x1 Soares NSF F8 9W OT PA P S1 Homo septens cDNA clone IMAGE:2148389 3'	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	xx10b04.x1 NCI_CGAP_Pan1 Homo sepiens cDNA clorie IMAGE:2837071 3' similar to gb:L20868 60S RIBOSOMAL PROTEIN L4 (HUMAN);
Top Hit Database Source	EST_HUMAN EST	EST_HUMAN 600	Hon	NT Hom	Hun tran	ISSPROT	SWISSPROT BO	EST_HUMAN we9	EST_HUMAN we9	SWISSPROT MUI	NT Han	Han	SWISSPROT COL	N For			EST_HUMAN Jy30	he3	EST_HUMAN qe8	EST_HUMAN qe6	SWISSPROT URI	EST_HUMAN QVC	EST_HUMAN QV3	SWISSPROT MY	EST HUMAN G12	Т	Т		EST_HUMAN RIB
Top Hit Acession No.	AA300640.1	BE299888.1	AL163247.2	AL163247.2	D13391.1		P98063	AI798902.1	AI798902.1	P08183	AJ271736.1	AL163207.2	P02461	U79410.1			R09942.1	AW468468.1	AI185642.1 E	AI185642.1	Q16831 S	BE062744.1	AW996538.1	Q28824 S	AA453647 1	T		4758977 NT	AW316976.1 E
Most Similar (Top) Hit BLAST E Value	7	7		2.0E-17	2.0E-17	1		2.0E-17	2.0E-17	1.0E-17	1.0E-17	1.0E-17	1.0E-17	1.0E-17	$\overline{}$	1.0E-17	١	1.0E-17	1.0E-17	1.0E-17	1.0E-17	1	1.0E-17	1.0E-17	1 0E-17	_	9.0E-18	8.0E-18	7.0E-18
Expression Signal	-	2.81	3.22	3.22	4.82	0.73	0.73	0.49	0.49	3.66	1.26	2.73	2.35	2.06		1.3	7.37	0.69	2.04	2.04	0.93	1.33	0.88	2.09	2 47	0.95	3.31	1.52	16.92
ORF SEQ ID NO:		32928		35658	36025				36170	26164		27204	27571	27795					32327	32328	32730	34324		37304	37649	27932	-	29156	25776
Exan SEQ ID NO:	21049		1		22807	L_	22926		22954	13507	14446	14503	14840	15059			16858	19136	19320	19320	19682	21182	22556	24000	24325	15192	22060	16518	13140
Probe SEQ ID NO:	8356	9769	9804	9804	10159	10278	10278	10307	10307	733	1703	1761	2109	2335		3554	4116	6366	6555	6555	6869	8490	2066	11394	11732	2474	8688	3766	339

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	Top Hit Descriptor	xx10b04.x1 NCJ_CGAP_Pan1 Homo sepiens cDNA clone IMAGE:2837071 3' similar to gb:L20868 60S RIBOSOMAL PROTEIN L4 (HUMAN);	RC3-OT0091-170300-011-d03 OT0091 Homo sapiens cDNA	xx10b04.x1 NCI_CGAP_Par1 Homo sapiens cDNA clone IMAGE:2837071 3' similar to gb:L20868 603 RIROSOMAL PROTEIN 14 (HLIMAN):	xx10b04.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE::2837071 3' similar to gb:L20868 60S RIBOSOMAI PROTFIN I 4 (HLIMAN)	Rattus norvegicus partial Gdn/Pn-1 gene for gila-derived nexin/protease nexin I, enhancer region	PROTEIN-GLUTAMINE GAMMA-GLUTAMYLTRANSFERASE (TISSUE TRANSGLUTAMINASE) (TGASE C) (TGC)	Homo saplens similar to high-mobility group (nonhistone chromosomal) protein 4 (H. saplens) (LOC63449), mRNA	Homo saplens chromosome 21 segment HS21C010	Homo saplens chromosome 21 segment HS21C048	H.sapiens DWA, DWB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING9, 9, 13 and 14 genes	Human aconitate hydratase (ACO2) gene, exon 4	qm65g11x1 Soares, placenta, 8to9weeks, 2NbHP8to9W Homo saplens cDNA clone IMAGE:1893668 3' similar to contains Alu repetitive element,	HUM411F05B Ciontech human fetal brain polyA+ mRNA (#8535) Homo sapiens cDNA cione GEN-411F05 5	Human endogenous retrovirus HERV-P-T47D	MRG-HT0161-221099-002-c06 HT0161 Homo saplens cDNA	Homo sapiens lymphocyte activation-associated protein (LOC51088), mRNA	Homo saplens lymphocyte activation-associated protein (LOC\$1088), mRNA	MR1-SN0035-060400-001-g11 SN0035 Homo sapiens cDNA	AV650547 GLC Homo sapiens cDNA clone GLCCGA02 3'	ho36h04.x1 NCI_CGAP_Ut1 Homo septens cDNA clone IMAGE:3039511 3' similar to contains MER29.b3	MER29 repetitive element ;	ho38h04.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:3039611 3' similar to contains MER29.b3 MER29 repetitive element;	nq24f1.s1 NCI_CGAP_Co10 Homo sapiens cDNA clone IMAGE:1144645 3' similar to gb:M26326 KFRATIN TYPE I CYTOSKEI FTAL 18 (HIMAN)	WISHARY NOT CRAP CARE Home seniors child clare MARCE 2202005 2	אוניטוויסיאן ואין בער בער בער איני איני איני איני איני איני איני אינ
	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	FST HUMAN	Z-L	SWISSPROT	LV	Į,	ZZ	N TN	NT	EST_HUMAN	EST_HUMAN	LN LN	EST_HUMAN	NT	TN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST HEIMAN	EST HIMAN	EST. LINIO.
	Top Hit Acession No.	7.0E-18 AW316976.1	7.0E-18 AW887542.1	8 AW316976 1	8 AW316976 1	8 X71791.2	8 P52181	11428155INT	8 AL 163210.2	8 AL 163246.2	8 X87344.1	8 U87929.1	8 A(280214.1	8 D61517.1	5.0E-18 AF087913.1	5.0E-18 BE143312.1	10242378 NT	10242378 NT	8 AW867182.1	B AV850547.1		4.0E-18 BE044076.1	BE044076.1	A A R 21 R 14 1	A 1728507 1	11100004.1
	Most Similar (Top) Hit BLAST E Value	7.0E-18	7.0E-18	7.01	7.015	6.0E-18	6.0E-18	6.0E-18	6.0E-18	6.0E-18	6.0E-18	6.0E-18	5.0E-18	5.0E-18	5.0E-18	5.0E-18	5.0E-18	5.0E-18	5.0E-18	5.05-18		4.0E-18	4.0E-18	4 OF-18	4.0E-187	7.01-101
	Expression Signal	16.92	1.33	3 41	3.41	1	3.02	284	0.72	1.61	1.74	3.29	21.7	0.98	1.2	6.25	3.47	3.47	3.4	4.18		1.37	1.37	4	7	1.14
	ORF SEQ ID NO:		33100	25778					34077		37260		28541	30384	30622		36828	38829				25580	25581			_
	SEQ ID NO:	13140	20024	13140	1	1	L	20842	ļ	23749	1	Į.	1	17768	17899	21312	23579	23579	24770	24978		12939	12939	44454	44404	2
	Probe SEQ (D NO:	339	7343	12492	12492	3289	4698	8148	8246	11079	11300	12241	1125	5047	5191	8620	10899	10899	12368	12895		12	121	1,2	1007	700

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<u> </u>	Top Hit Detabase Source Source Source Source Source EST_HUMAN EST_HUMAN EST_HUMAN NT NT NT NT NT NT EST_HUMAN EST_HU	Top Hit Acess No. No. No. Ho. Ho. Ho. Ho. Ho. Ho. Ho. Ho. Ho. H	Most Simi (Top) High BLAST I BLAST I Sold BLAST I Sold Blast I Sold Bl	Sign Sign	36728 35788 35788 35789 37180 31180 31181 32128 34171 34171 34419 34419 34419 34420 37016	 	Probe SEQ ID NO: 10894 11736 1
EST38/00/ MAGE resequences, MAGN Homo saplens cDNA	SWISSPROT	P08548	8.0E-19	104		17110	4372
Г	EST HUMAN	AW974902.1	8.0E-19	1.25		13786	1026
	ST_HUMAN	AA281961.1	9.0E-19	1.88	25952	13316	11901
THINDS A NO! COAD GORI Home content along MACOF TARGET II.							
Homo saplens mRNA for KIAA1143 protein, partial cds	F	AB032969.1	9.0E-18	4.82	37016	23742	77011
Homo sapiens chromosome 21 segment HS21C003	L	AL163203.2	9.0E-18	2.54	34420	70717	000
Homo sepiens chromosame 21 segment HS21C003	Į.	AL163203.2	9.0E-12	5,	100	2007	3 6
Т	ES! HOMAN	FU8086.1	9.0E-18	4.4.	24440	24.267	B.C.B.
1	TO L HOMAIN	AAZO1801.1	9.VE-16	74.47		20443	7747
	MANUEL FOR	A A 28 10 64 1	0 OF-40	3 24	25952	13316	88
	EST_HUMAN	AA281961.1	9.0E-18	5.34	25952	13316	632
Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	Ė	AF003529.1	1.0E-1	4.23	31082	1	812
(TLA-T) gene, Roket gene, and sodium phosphate transporter (NPT3) gene, complete cds	- 1	001020.1	100	100	34000	1	12430
Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis		. 191328 1	1.0F-1	4.22	36653		9789
	EST HUMAN	AI148288.1	1.0E-1	1.43	34171	21034	8341
0269409 x1 Spatres settlement filmwharle NNHSE Dame conjunction (NAS)							
Homo sapiens chromosome 21 segment HS21C080	F	JAL 163280.2	1.0E-1	1.63	32128	-	6363
Homo sepiens mRNA for Na,K-ATPase alpha-subunit, complete cds	NT	3 D00099.1	1.0E-1	2.94		┙	5483
Homo sapiens mRNA for Na,K-ATPase alpha-subunit, complete cds	ΙΝ	3 D00099.1	1.0E-1	2.94	ı		2483
	EST_HUMAN	5.1	1.0E-1	3.63	30707		5271
	EST_HUMAN	8 T95406.1	1.0E-1	1.02			4382
П	EST_HUMAN	8 BE256097.1	2.0E-	1.45		- 1	12174
	EST_HUMAN	8 AW 151299.1	2.0E-	3.91			11736
T		10001					
	TOT LINAAN	8 AM(470704 4	2.0E-4	~			10894
	EST_HUMAN	8 AW 151673.1		1.66		i	8947
	EST_HUMAN	8 AW151673.1	-			_ 1	9947
x(87ed or NC) CCAP Good Home continue about a late of contract of the contract							
	Source	•	Value		<u></u>	Ö	Ö
		à .	Most Simile (Top) Hit	Expression			Probe SEQ ID
robes Expressed in Brain	gie Exon P	un l					
mhae Evnressad in Brain	idle Exon P						

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	Top Hit Descriptor	Homo sapiens DEAD/H (Asp-Gtu-Ala-Asp/His) box polypeptide 6 (RNA helicase, 54kD) (DDX6) mRNA	Rattus norvegicus cp151 mRNA, partial cds	BETA CRYSTALLIN A2	tb01c08.x1 NCI_CGAP_Lu26 Homo saplens cDNA clone IMAGE:2052302.3"	zi60b01.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:4351453'	PM0-CT0248-131099-001-g01 CT0248 Home sapiens cDNA	OLFACTORY RECEPTOR 6 (M50)	OLFACTORY RECEPTOR 6 (M50)	Homo sapiens Xq pseudoautosomal region; segment 1/2	DKFZp762F192_r1 762 (synonym: hmel2) Hamo saplens cDNA clane DKFZp762F192 5'	ZONA PELLUCIDA SPERM-BINDING PROTEIN B PRECURSOR (ZONA PELLUCIDA GLYCOPROTEIN 2P-X) (RC55)	Homo sapiens partial IL-12RB1 gene for IL-12 receptor beta1 chain, exon 14	x/87b02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone iMAGE:2684171 3' similar to contains	element MSR1 repetitive element ;	Homo sapiens mRNA, chromosome 1 specific transcript KIAA0501	602130910F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4287674 5	Homo sapiens mannosidase, beta A, Iysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (I IBE2D3) กลคละ ความกลุ่ยล cde	BETA-2 ADRENERGIC RECEPTOR	BETA-2 ADRENERGIC RECEPTOR	LIM-ONLY PROTEIN 6 (TRIPLE LIM DOMAIN PROTEIN 6)	LIM-ONLY PROTEIN 6 (TRIPLE LIM DOMAIN PROTEIN 6)	AV708136 ADC Homo sapiens cDNA clone ADCAMA11 5	Homo sapiens NPD008 protein (NPD008) mRNA, complete cds	Homo sapiens similar to aldo-keto reductase family 1, member B11 (aldose reductase-like) (H. sapiens)	(LOC63222), mRNA	M.musculus mRNA for TPCR33 protein	Homo saplens phorbolin I protein (PBI) mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C001	qo91e02x1 NCI_CGAP_Kid5 Homo saplens cDNA clone IMAGE:1915898 3' similar to TR:Q68386 Q69386 POL/ENV GENE;	AV731382 HTF Homo sapiens cDNA clone HTFAZC06 5
0: 1: IOV 0: 0: 0: 0: 0: 0: 0: 0: 0: 0: 0: 0: 0:	Top Hit Database Source	⊢ N	NT.	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	Į,	EST_HUMAN	SWISSPROT	TN		EST_HUMAN	L	EST_HUMAN	LN.	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	EST_HUMAN	IN	1	LN	NT	TN	LN	EST_HUMAN	EST_HUMAN
	Top Hit Acession No.	4758139 NT	9 AF092090.1	9 P26444	9 Al344951.1	9 AA705684.1	19 AW852930.1	P34986	P34986	AJ271735.1	19 AL120817.1		19 AJ297699.1		9 AW183725.1	9 AB007970.1	4.0E-19 BF697362.1	Q A F 224669 1	9 028997	9 0 28997	9 043900	9 043900	9 AV708136.1	9 AF223467.1		11432214 NT	9 X89685.1	9 AF165520.1	9 AL163201.2	9 Al311783.1	9 AV731382.1
	Most Similar (Top) Hit BLAST E Value	7.0E-19	7.0E-19	7.0E-19	7.0E-19	7.0E-1	6.0E-1	6.0E-19 P34986	6.0E-19 P34986	6.0E-19	6.0E-19	5.0E-19	6.0E-19		5.0E-19	4.0E-19	4.0E-19	4 0E-19	3.0E-19	3.0E-19	3.0E-19	3.0E-19	3.0E-19	3.0E-19		3.0E-19	3.0E-19	3.0E-19	2.0E-19	2.0E-19	2.0E-19
	Expression Signal	1.51	2.34	6.0	0.51	2.05	1.34	1.36	1.36	1.3	1.04	5.36	1.03		7.45	1.68	1.02	70.0	1.04	1.04	66.0	66.0	1.12	0.64		2.79	1.15	23.34	21.33	1.03	0.57
	ORF SEQ ID NO:	27708	32129	32969					96262		30301	31485	36191		١	25956	28136	30773		29220		29623					33548		28017		31706
	Exan SEQ ID NO:	14970									17692	18558	22971	l	- 1	13324	15398	18115	1	I.		Ŀ	17150	18006		ı	20430	24709	15279	17148	1 1
	Probe SEQ ID NO:	2242	6364	7199	9911	12036	3761	4430	4430	4747	4967	2929	10324		11625	2	2689	5311	3833	3833	4253	4263	4413	5198		7283	8359	12264	2565	4411	5963

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SEQ ID SEQ ID NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:	ORF SEQ ID NO: 28949 28605 33763 33668 33689 33625 34673	Express Sign	Most (10 kg)	12633	Top Hit Database Source EST HUMAN NT SWISSPROT EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN	Top Hit Descriptor Ind6c04.s1 NCI_CGAP_Pr4 Homo septens cDNA clone IMAGE:1043718 similar to contains MER29.b2 MER29 repetitive element; Homo septens ribosomal protein L13a (RPL13A), mRNA ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY 501441231F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:39162316; AV725123 HTC Homo sapiens cDNA clone HTCBTA01 5 AV725123 HTC Homo sapiens cDNA clone HTCBTA01 5 AF075301 Human fetal liver_spleen_1NFLS_51 Homo sapiens cDNA clone IMAGE:418191 3' similar to contains MER30:11 MER30 repetitive element; 2n78d08:1 Soares_fetal_liver_spleen_1NFLS_51 Homo sapiens cDNA clone IMAGE:418191 3' similar to contains MER30:11 MER30 repetitive element; 2n78d08:1 Soares_fetal_liver_spleen_1NFLS_51 Homo sapiens cDNA MR3-HT0487-15020-0113-g01 HT0487 Homo sapiens cDNA Mus musculus MMAN-g mRNA, complete cds Mus musculus MMAN-g mRNA, complete cds
11111		0.84 1.34 5.15 1.33			SWISSPROT NT SWISSPROT EST_HUMAN EST_HUMAN	HYPOTHETICAL PROTEIN DJ845024.1 Homo saplens chromosome 21 segment HS21C047 HISTONE H2B C (H2B/C) E264g03.x1 NCI_CGAP_Ov35 Homo saplens cDNA clone IMAGE:2283396.3' QV3-DT0043-090200-08004 DT0043 Homo saplens cDNA Human BXP21 cene
2135 14895 4185 16926 4582 17317 8833 21525 10219 22867	29557 29557 29944 36078		3.0E-20 3.0E-20 3.0E-20	3.0E-20 U03888.1 3.0E-20 P23273 3.0E-20 A4037618.1 3.0E-20 D14547.1 3.0E-20 BF185264.1	SWISSPROT EST_HUMAN NT EST_HUMAN	Human BXP21 gene OLFACTORY RECEPTOR-LIKE PROTEIN 114 A286b12.s1 Scares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:484895 3' similar to contains L1.t3 L1 repetitive element; Human DNA, SINE repetitive element 601843591F1 NIH_MGC_54 Homo sapiens cDNA done IMAGE:4064343 5' RETROVINGUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE;
<u> </u>	37408 37409 31118	1.5 1.5 2.86 3.12		4.1 22.1 888.1		groodozxi NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1864803 3' similar to contains Alu repetitive element: agrodozxi NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1864803 3' similar to contains Alu repetitive element: element: 601614180F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3915522 5' x24e10.x1 NCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:2761098 3' similar to SW:RS5_MOUSE P97461 40S RIBOSOMAL PROTEIN S5.;

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Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21C018	Homo sapiens dNT-2 gene for mitochondrial 5(3')-deoxyribonucleotidase (dNT-2 gene), exons 1-5	Human chromosomal protain HMG1 related gene	RC0-CT0301-271199-031-F03 CT0301 Homo sapiens cDNA	zg/3403.s1 Soares. fetal, heart. NbHH19W Homo sapiens cDNA clone IMAGE:399981 3' similar to gb:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN);contains THR.t3 OFR repetitive element;	Homo saplens PTD013 protein (PTD013), mRNA	601304125F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638310 5	PM1-HT0454-080100-002-h09 HT0454 Homo sapiens cDNA	Homo saplens protein tyrosine phosphatase, non-receptor type 21 (PTPN21), mRNA	601649871F1 NIH_MGC_74 Hamo saplens cDNA clane IMAGE:3833880 57	Homo sapiens melanoma antigen, family C, 1 (MAGEC1), mRNA	he05e10.x1 NCI_CGAP_CML1 Homo saplens cDNA clone IMAGE: 2918154 3'	783411.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:3303573 3' similar to contains OFR.t1 OFR repetitive element :	ZINC FINGER PROTEIN GLI1 (GL-1)	ZINC FINGER PROTEIN GLI1 (GLL1)	zi72c04.r1 Soares_testis_NHT Homo sapiens cDNA done IMAGE:727878 5'	0088e08.s1 NCI_CGAP_Kid5 Homo saplens cDNA clone IMAGE:1573094 3' similar to TR:Q16530 Q16530 PMS3 MRNA ;contains OFR.t1 OFR repetitive element;	Rattus novegicus mRNA for rTIM, complete cds	Human hereditary haemochromatosis region, histone 2A-lika protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	Hamo sapiens chromosome 21 segment HS21 C002	zq15d06.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:629771 3'	Homo sapiens chromosome 21 segment HS21C001	Homo saplens LGMD2B gene	Homo sapiens dNT-2 gene for mitochondrial 5′(3′)-deoxyribonucleotidase (dNT-2 gene), exons 1-5	Homo sapiens dNT-2 gene for mitochondrial 5'(3')-deoxynbonucleotidase (dNT-2 gene), exons 1-5
Top Hit Database Source	NT	FZ	N	EST_HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	IN	EST_HUMAN	EST HUMAN	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	N	NT	NT	EST_HUMAN	NT	NT	NT	TN
Top Hit Acession No.	AL163218.2	AJ277557.1	D14718.1	AW856922.1	AA723404.1	7706668 NT	BE408611.1	BE162737.1	5902031 NT	BE968839.1	4885474 NT	AW440864.1	BE856505.1	Q91690	Q91690	AA393574.1	AA970713.1	AB019576.1	U91328.1	AL163202.2	AA218891.1	AL163201.2	AJ007973.1	AJ277557.1	AJ277557.1
Most Similar (Top) Hit BLAST E Value	7.0E-21	7.0E-21	7.0E-21	7.0E-21	7.0E-21		6.0E-21		5.0E-21		5.0E-21	5.0E-21	5.0E-21		5.0E-21		4.0E-21				3.0E-21		3.0E-21	3.0E-21	3.0E-21
Expression Signal	0.79	1.53	10.76	0.86	2.19	1.75	0.83	0.6	7.0	2.91	5.58	6.0	0.86	0.44	0.44	2.83	1.81	3.27	0.63	2.0	0.94	1.24	4.31	0.68	0.68
ORF SEQ ID NO:	32100	34121	34407		36525	37184	29454		26334	29697	30112		32700	38349	36350		27168	32544	35527	35552	27280	27736	28485	30919	30920
Exon SEQ ID NO:	19110	20981	21268	['	23288	23897	16827	21724	13670		17481	19582	19654		23120	1	14469	19518		22356	Ь.	14998	15843	18211	18211
Probe SEQ ID NO:	6340	8287	8576	10013	10594	11234	4083	9034	506	4330	4749	6865	2 H69	10474	10474	11986	1727	6772	0896	9705	1829	2272	3078	5412	5412

Page 244 of 536 Table 4

ts30f03.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2230109 3' sImilar to TR:Q99884 Q99884 HYPOTHETICAL 51.1 KD PROTEIN; hi09g01 x1 NCI_CGAP_Kld13 Homo sapiens cDNA clone IMAGE:3148266 3' similar to contains MER29.b3 Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA 1294e03 x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2298204 3' similar to TR:Q15408 Q15400 NEUTRAL PROTEASE LARGE SUBUNIT; ar88d12.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:2162343 3'
DKFZp43410830_11 434 (synonym: hiss3) Homo sapiens cDNA clone DKFZp43410830 5'
9947e05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1838338 3' similar to gb:M84241 QM
PROTEIN (HUMAN); 601680636F1 NIH_MGC_83 Homo saplens cDNA clone IMACE:3951008 5'
601680636F1 NIH_MGC_83 Homo saplens cDNA clone IMACE:3951008 6'
Homo saplens putative 8-hydroxyguenine DNA glycosylase gene, complete cds

NAGCO4.s1 NCI_CGAP_P14 Homo saplens cDNA clone IMAGE:1043718 similar to contains MER29.b2 ze97a12.r1 Soares_fetal_heart_NbHH19W Homo saplens cDNA clone IMAGE:366910 57 ze97a12.r1 Soares_fetal_heart_NbHH19W Homo saplens cDNA clone IMAGE:366910 57 601844485F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4084945 6 QV0-HT0103-091199-050-g11 HT0103 Homo sapiens cDNA AU136779 PLACE1 Homo sapiens cDNA clone PLACE1005052 6' Top Hit Descriptor CM1-NN0063-280400-203-h08 NN0063 Homo sapiens cDNA Homo sapiens chromosome 21 segment HS21C013 QV3-HT0458-170200-090-912 HT0458 Homo sapiens cDNA RC4-BT0311-141199-011-h06 BT0311 Homo sepiens cDNA RC1-0T0083-100800-019-908 OT0083 Homo saplens cDN AV661044 GLC Homo sapiens cDNA clone GLCGOA10 3 Homo sepiens mRNA for KIAA0397 protein, partial cds Homo sapiens mRNA for KIAA0397 protein, partial cds Homo sepiens chromosome 21 segment HS21C001 Homo sepiens chromosome 21 segment HS21C001 Homo sapiens hyperion gene, exons 1-50 ZONADHESIN PRECURSOR ZONADHESIN PRECURSOR Single Exon Probes Expressed in Brain MER29 repetitive element; MER29 repetitive elemen EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN Top Hit Database Source HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HCMAN EST HUMAN EST_HUMAN EST_HUMAN NT SWISSPROT EST_HUMAN EST HUMAN HUMAN EST_HUMAN EST EST Top Hit Acession 5730038 3.0E-21 AL163213.2 2.0E-21 BE163247.1 2.0E-21 AB007857.2 2.0E-21 AB007857.2 3.0E-21 AV661044.1 3.0E-21 BF184739.1 3.0E-21 BF361093.1 AW897760.1 BF184739.1 2.0E-21 BE350127.1 2.0E-21 BE973829.1 2.0E-21 BE973829.1 2.0E-21 AF176815.1 9.0E-22 AI702438.1 9.0E-22 AL163201.2 9.0E-22 AL163201.2 2.0E-21 AA027211.1 2.0E-21 AA027211.1 BE141785.1 AU136779.1 BE064410.1 2.0E-21 AJ010770. AA557657.1 1.0E-21 AI601264.1 1.0E-21 AL079752.1 2.0E-21 AI624582. 1.0E-21 AI223104.1 1.0E-21 57 3.0E-21 (Top) Hit BLAST E 1.0E-21 2.88 0.65 1.98 8. 1.85 0.68 0.5 1.55 Expression 1.3 9.87 9. 9.0 1.27 2.62 Signal ORF SEQ 35429 26342 30714 26343 28098 28099 30890 31184 31185 33996 34557 37247 ÖΝΟ 26652 34336 32847 29747 SEQ ID 18864 18447 19451 15354 15354 13681 13944 18196 23865 23950 23950 18288 20864 20955 21414 13982 14128 ġ 19781 17114 24712 19165 23130 21194 21194 SEQ ID 5852 6086 9592 6969 12533 <u>+</u> 914 1192 2844 5396 914 5489 8170 8261 11289 10991 12272 138 138 138 138 138 7092 10484 8502 4377 8502

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Probe SEQ ID NO:		S O	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian No.	Top Hit Database Source	Top Hit Descriptor
10691	23382			9.0E-22	22 AV761874.1	EST_HUMAN	AV761874 MDS Hamo sapiens cDNA clone MDSCCG05 5'
11707	Ш	37627	1.34	9.0E-22	22 AU140358.1	EST_HUMAN	AU140368 PLACE2 Homo sapiens cDNA clone PLACE2000394 5'
928			59.5	8.0E-22	2 BE144748.1	EST_HUMAN	CM0-HT0179-281099-076-h05 HT0179 Homo sapiens cDNA
797	20492		3.72	8.0E-22	22 AA046502.1	EST_HUMAN	zk87a08.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:487858 5'
650				7.0E-2	2 AL163246.2	Z	Homo sapiens chromosome 21 segment HS21C046
4250		29616	2.21			SWISSPROT	ALPHA-2-MACROGLOBULIN PRECURSOR (ALPHA2M)
4977	17700					ΙN	Homo sapiens gene for activin receptor type IIB, complete cds
8590	21282					N	Homo sapiens HSPC220 mRNA, complete cds
8731	21423	34568			7.0E-22 M78590.1	EST_HUMAN	EST00738 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA done HFBCF07
9502	22155				7.0E-22 AF009660.1	NT	Homo sapiens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
4038	16783	29413	0.98		2 AA405040.1	EST_HUMAN	zu65d10.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:742867 51
8140	20834				6.0E-22 AW029123.1	EST_HUMAN	wx05g07.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2542812 3'
6424	19192	32188	3.76		AL163303.2	INT	Homo sapiens chromosome 21 segment HS21C103
10217	22865				5.0E-22 U60822.1	NT	Human dystrophin (DMD) gene, exons 7, 8 and 9, and partial cds
							naa27b06.x1 NC_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:3255898 3' sImilar to contains Alu
12499	24854		2.22	Į	5.0E-22 BF476511.1	EST_HUMAN	repetitive element;
3627	16380		0.85		AJ271735.1	F	Homo saplens Xq pseudoautosomal region; segment 1/2
8004	20699	33827	0.45		AV703223.1	EST_HUMAN	AV703223 ADB Homo sapiens cDNA clone ADBAUE12 6
8312	25428		3.11	4.0E-22	AL163202.2	NT	Homo sapiens chromosome 21 segment HS21C002
10623	23316	36556	2.47	4.0E-22	4.0E-22 BF218030.1	EST_HUMAN	601882813F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:4095434 5'
12657	24957		2.08	4.0E-22		LN	Homo sapiens chromosome 21 segment HS21C009
939	13706		1.58	3.0E-22	3.0E-22 AI469679.1	EST_HUMAN	tm14h10.x1 NCI_CGAP_Co14 Homo saplens cDNA clone IMAGE:2156811 3' similar to gb:L19593 HIGH AFFINITY INTERLEUKIN-8 RECEPTOR B (HUMAN).contains L1.t1 L1 repetitive element :
						Γ	wi66b04.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2429839 3' stmilar to SW:RL21_HUMAN
2575	15289	28026	0.92	3.0E-22,	1	EST_HUMAN	P46778 60S RIBOSOMAL PROTEIN L21.;
3662	16415		1.46	3.0E-22	3.0E-22 D14718.1	L	Human chromosomal protein HMG1 related gene
4740	17400	2777	ď	000	7 300001	MANAGE TO TO T	qb28c07.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:1697580 3' similar to
2 2	2000		2,4	2000	0.0C-22 A1030 (20.1	Т	OVICE INTO A STATE OF THE STATE
87 0	2007		0.0	3.05-22	Z BE130613.1	T	CVU-H 10386-CUCZOU-099-112 H 10368 Homo sapiens cDNA
813 4	20828		2.48	3.0E-22	-	T_HUMAN	RC5-BT0707-150300-021-H10 BT0707 Homo sepiens cDNA
8228	20952		0.97	3.0E-22			R. rattus RY2G5 mRNA for a potential ligand-binding protein
8258	20952	34089	0.97	3.0E-22 X60660.1			R.rattus RY2G5 mRNA for a potential ligand-binding protein
1946	14681		2.29	2.0E-22 N24942.1			yx73d05.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:267369.31
2528	15242	27981	2.15	2.0E-22	24916	SWISSPROT	IMMEDIATE EARLY GENE 13 PROTEIN PRECURSOR

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		_		TT	T	7	Т	-	_	Т	Т	Т	Т	Т	_	_	T	_	_	_	_	_	_							-
Single Exon Probes Expressed in Brain	Top Hit Descriptor		Hono saplens protein kinase. AMP-achireled	PM1-ST0282-261199-001-412 ST0282 Homo saplens cDNA	9b.X72308 MONOCYTE CHEMOTACTIC PROTECTION 2 PROTECTION 4 clone IMAGE:322873 6' similar to	RCD-TN0079-150900-025-h12 TN0079 Hamo saplens cDNA	MER29.13 MER29 reportition along a plens cDNA clone IMAGE:1878289 3' similar to containe	NV4h11.s1 NCI CGAP Pro Home	nv04h11.s1 NCI_CGAP_Pr22 Homo seniers cDNA clone IMAGE:1219289 3'	ha24f04.x1 NCI_CGAP_Kid12 Home sanions contact in the contact in t	Homo saplens chromosome 21 seament HS21Cnsp	PM4-SN0020-010400-009-h02 SN0020 Home sastore - DN14	Human familial Alzhetmer's disease (STM2) nane com-Li	Human DNA, SINE repetitive element	MR0-BT0659-220200-002-h07 BT0689 Home and a second and a second a	9209b07.x1 NCI_CGAP_CIL1 Homo septems cDNA clara IMA CE.	ER29 repetitive element; D9b07 vd NCI CCAD OF CONTRIBUTE CONTRIBUTE OF CONTRIBUTE MER29.b2	MER29 repetitive element	L2-UM0076-070400-084-E11 I IMAGG L	Gallus gallus Dach2 protein (Dach2) w Data	AV647246 GLC Home sapiens child also of Street	Homo saplens Not56 (D. melanopaster) like profits Angered	Reftus norvegicus RIM1B (Rim1B) mRNA complete etc.	Hano sepiens chranosame 21 segment HS210049	(UBE2D3) genes, complete cds	Homo saplens mannosidase, beta A, Ivsosomal (MANRA) rene	(UBEZD3) genes, camplete cds	SW:MV10_MOUSE P23249 PROTEIN MOV.16 .	Homo sepiens chromosome Xq28 melanoma antigen family A2a (MAGEA2A), melanoma antigen family A12 (MAGEA12), melanoma antigen family A2b (MAGEA2B), melanoma antigen family A3 (MAGEA2B), melanoma antigen family A3 (MAGEA2B), melanoma antigen family A3 (MAGEA3B),	(WASCEAS), cattacting and LI>
gie Exon Pro	Top Hit Database Source			EST_HUMAN		EST_HUMAN		Г	П	HUMAN	П	HOMAN		\neg	EST HUMAN N		NAMON NAMON NA		T HUMAN	Ť	T_HUMAN			운 운		Ÿ	(0)	EST HUMAN SW	<u> </u>	
	Top Hit Acession No.		2 0E-22 AM 84-12	AW817794.1	-22 W39456.1	T	2.0E-22 AI276522.1		22 AA715315.1	1	22 AN BREE17 4	T	T	22 DE004007	1	22 Al365435 1						31852		N 7.84-70	AF224669.1 NT	AF224669 1 NT		AI209130.1 EST	U82671.2 NT	
	Most Similar (Top) Hit BLAST E Value	200	20E-22	77-20-7	2.0E-22		20E-22/	Z.OE-22/	2.0E-22	2 OF 22 A	1.0F-22	1 0F-2211	1.0F.22.D	1 OF 22 B		1.0E-22 AI		1.0E-22 Al365435.1	9.0E-23 AW 802801.1	7 0E 22 A1/2 (E.C.)	7.0E-23/AV64/246.1	R 0F-23 A E400000	6.0E-23 At 489340.0	2	6.0E-23 AF	6.0E-23 AF2		8.0E-23 AI20	6.0E-23 U82	
	Expression Signal	4 41	1.17	,	3.58	,	3 8	9 6	188	3.71	1.79	1.7	1.37	0.89		0.79	- G	5 RO	02.0	2.65	4 18	1.72	1.39	;	2	1.5		3.28	4.09	
	ORF SEQ ID NO:		29567	1	Ц	35440		35541	37884	30983	27320	28038	28814	33416		36328	36320	+	28959		36895		29601	31105	-	31106	10000	2000	30844	
	SEQ ID NO:	4 16172	16941	25075	П	22255		22347		_1	14609	15302	16163	20308		23097	23097	24984	16312	16065	23642	16184	16976	24540	-	24540	24663	-	18160	
	Probe SEQ ID NO:	3414	4200	5761	6084	8602	9696	9696	11761	11872	1871	880	3405	384		10451	10451	12704	3557	3305	10966	3427	4236	12005		12005	12192	1	5358	
																			_						Ц_	ㅗ		Щ.		

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Most Similar (Top) Hit BLASTE No. Source
23 AF179818.1 NT
5.0E-23 AF179818.1 NT Pongo pygmaeus olfactory receptor (PPY116) gene, partial cds
23 AL 163227.2 NT
3.0E-23 AL163227.2 NT Homo sapiens chromosome 21 segment HS21C027
2/35g09_r1 Scares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:503968 5' similar to
TIME TO THE
2 270004.1 IN
NT
3.0E-23 AW897927.1 EST_HUMAN RC3-NN0066-270400-011-h01 NN0066 Hamo sapiens cDNA
2.0E-23 AJ289880.1 NT Homo saplens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene
3 M55270.1 NT
2.0E-23 P22105 SWISSPROT TENASCIN-X PRECURSOR (TN-X) (HEXABRACHION-LIKE)
23 P22105 SWISSPROT
S AI201458.1 EST HUMAN
2.0E-23 BE165980.1 EST_HUMAN MR3-HT0487-150200-113-g01 HT0487 Homo sapiens cDNA
3 H59931.1 EST_HUMAN
E-23 H59931.1 EST_HUMAN yr16a02.r1 Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:205418 5'
4 (CYP344) and cytochrome P450 polypeptide 7 (CYP347) genes, complete cds; and cytochrome P450 polypeptide 7 (CYP347) genes, complete cds; and cytochrome P450 polypeptide 7 (CYP345) gene partial cds
3 AL163303.2 NT
Ā
2.0E-23 AF009660.1 NT Homo saplens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
1.0E-23 AL163252.2 NT Homo saplens chromosome 21 segment HS21C052
3 AL163210.2
1.0E-23 BE378471.1 EST_HUMAN 601236456F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3608653 S'
Zv82c06.r1 Soares, testis, NHT Homo sapiens cDNA clone IMAGE:782698 5' similar to contains PTR5.t2
3 AA448097.1 EST_HUMAN
3 BE409643.1 EST_HUMAN
1.0E-23 BE409643.1 EST_HUMAN 601301762F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3636254 5'
HUMAN
E-24 11422027 NT Homo sapiens capping protein (actin filament) muscle Z-ltne, alpha 2 (CAPZA2), mRNA

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					5	IGIO EXOTI PIO	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3853			1.49	7.0E-24	7.0E-24 AW937954.1	EST HUMAN	10V0-DT0047-170200-122-ang PT0047 Home and and and and and and and and and and
5087	17806		0.95	7.0E-24	-24 AL039498.1	EST HUMAN	DKFZ0434A2311 11 434 (singnum hises) Home content DNIA - Long Story - 1011 21
10536	23233		1.33	7.0E-24	24 AW303317.1	EST HUMAN	xv1703.x1 Soares_NFL_TGBC_S1 Homo septents CDNA clone IMAGE:2813403 3' similar to contains Alu repetitive element contains. MERA to MERA to MERA to MERA to Septents.
690			2.72	6.0E-24	24 AB001421.1	LN	Macaca fuscale mRNA for Teetle, Specific Ductor VTSBN
818		26256	11.74	6.0E-24	6.0E-24 AL163249.2	Į.	Homo sapiens chromosome 21 segment HS210040
3953	16703	. 28342	7.9	5.0E-24	24 AJ229043.1	Ę	Homo sapiens 959 kb contid between AMI 1 and CBR1 or characters 224.22
7657	20321	33430	0.58	5.0E-24	24 AF223391.1	LZ	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial ode, alternatively spliced
11595	24194	37513	1.45	5.0E-24	24 AW514229.1	EST_HUMAN	hd24b03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2910413 3' similar to TR:094851 C94851 KIAA0750 PROTEIN :
5840	18628	31563	3.85	4.0E-24	24 AA594178.1	EST HUMAN	nn31h05.s1 NCI_CGAP_Gas1 Homo saplens cDNA clone IMAGE:1085529 3' similar to SW:POL_MLVRK
8581	21273	34411	1,35	4.0E-24		EST HUMAN	RC3-ST0197-130100-014-808 ST0497 Home confirm
11133	ı	37078	1.95	4.0E-24 E	Γ	Т	601078812F1 NIH MGC 12 Homo senions china class (1978-1974 NIH MGC 12 Homo senions china class (1978-1974)
12361	ļ	31062	4.89	4.0E-24	Γ	Т	Homo sapiens mRNA for KIAA1003 protein partiel of
12595	24951	30986	1.77	4.0E-24	8318		Homo saplens G-2 and S-phase expressed 4 (CTSE4) - DNA
8322	21015		2.85	3.0E-24 A	3.0E-24 AW614871 1	Z	th68c08.x1 NCI_CGAP_GU1 Homo sepiens cDNA clone IMAGE:2867850 3' similar to contains MER29.b2
8377	21070		1.57	3.0E-24 A	T	Т	FST374149 MAGE recentance MAGG U
9365	21940	35114	4.33	3.0E-24 A	Γ	Т	Homo septiens chromosome 24 segment DC2400s
12438	li	31045	1.41	3.0E-24 B	Γ	T HUMAN	801810449F1 NIH MGC 46 Homo september China IMA CE 2008 81
23 28 28 28 28 28	- !	27806	2.72	2.0E-24 A	2.0E-24 AA167539.1	EST_HUMAN	zp11f09.r1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE: 606484 gi
3778	- 1		1.01	2.0E-24 A		EST_HUMAN	RC3-NN0068-090500-021-b03 NN0068 Homo sapiens cDNA
7374	- 1	33135	0.81	2.0E-24 A			Mus musculus rho/rac-interacting citron kinase (Crit) mRNA complete cds
7379	- 1	33138	0.65	2.0E-24 A	Γ	EST HUMAN	AJ003536 Selected chromosome 21 cDNA ilham Homo and home
8639	21331	34476	3.28	2.0E-24 A	2.0E-24 AL119158.1	Г	DKFZp761L1712_r1 761 (sworym: hamv2) Home sapiens cDNA close DKFZp7611 1712_r1 751
8676	21368		0.98	2 DE-24 H69214 1		TO LOUIS	y/92b09.r1 Scares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:212729 6' similar to contains
9754	22405	35611	76 0	2 OE-24 A	T	Ţ	MICACO I Special Control ()
9754	22405	35812	76.0	2 OE 24 A	T	Т	UT 809.X1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2138008 3'
11825	24409	37744	1 34	2000	T	Т	107 AUGUST NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2138008 3'
11825	24409	37745	2 6	2.0E-24 A	2.0E-24 AWGGGGGGG	Т	MR1-SN0063-040500-001-a06 SN0063 Homo sapiens cDNA
12281	26377		7 44	2 OF 24 M		ES - AUMAN	MK1-SN0063-040500-001-e06 SN0063 Homo sapiens cDNA
	1 1 1 1 1	1	1.41	4.VE-24 IV	1	ᅱ	Human O family dispersed repeat element

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acessian Na.	Top Hit Database Source	Top Hit Descriptor
1691	14435	27131	2.69	1.0E-24	7706340 NT	NT	Hamo sapiens CGI-127 protein (LOC51646), mRNA
2679	15388		1.63	1.0E-2	4 AW820194.1	EST_HUMAN	QV0-ST0294-100400-185-c10 ST0294 Homo saplens cDNA
3020		28433	1.49		1.0E-24 D86423.1	LΝ	Mus musculus mRNA for HGT keratin, partial cds
4237			1.71		4 AF143313.1	NT	Homo sapiens PTEN (PTEN) gene, excn 2
7447			4.32		1.0E-24 AL163303.2	NT	Hamo saplens chromosome 21 segment HS21C103
7630		33404	0.81		1.0E-24 BE144526.1	EST_HUMAN	MR0-HT0166-271199-005-d09 HT0166 Hamo sapiens cDNA
7845		33667	2.09		1.0E-24 AW801164.1	EST_HUMAN	CM0-NN1010-130300-281-d07 NN1010 Homo saplens cDNA
11699	24294	37619	1.31		7706707 NT	NT	Homo saplens putative secreted protein (SIG11), mRNA
							ne92e10.s1 NOI_CGAP_Kid1 Homo saplens cDNA clone IMAGE:911754 similar to contains MER1.b2
4939	17667	30275	2.33		7.0E-25 AA483944.1	EST_HUMAN	MER1 repetitive element ;
							ne06a09.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:880408 3' similar to contains THR.b2 THR
8117	20811	33946	6:29		7.0E-25 AA468646.1	EST_HUMAN	repetitive element;
11701	24286	37622	3.28		7.0E-25 AA583540.1	EST HUMAN	nr25h08.s1 NCI_CGAP_Pr1 Homo saplens cDNA clone IMAGE:914843 similar to SW:R14A_YEAST P36105 PROBABLE 60S RIBOSOMAL PROTEIN L14EA.
888	L		4.9		5 W87623.1	EST HUMAN	zh65h07.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:416989 5'
7622	20288	33397	8.34	6.0E-25	7305360 NT	LN TN	Mus musculus otogelin (Otog), mRNA
1647		27083	1.18	5.0E-25	5.0E-25 AW850271.1	EST_HUMAN	L3-CT0219-161199-031-D04 CT0219 Homo saplens cDNA
11286			2.44	5.0E-2	5 AW979107.1	EST_HUMAN	EST391217 MAGE resequences, MAGP Homo sapiens cDNA
1429	L	26861	2.25	4.0E-25	T98107.1	EST_HUMAN	ye56h04.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:121783 5
3397	16155		3.04		AW887671.1	HUMAN	PM3-OT0093-280200-001-g07 OT0093 Homo sapiens cDNA
4282	17021		2.83		4.0E-25 BE170957.1	EST_HUMAN	QV3-HT0543-140400-148-e11 HT0543 Homo sapiens cDNA
3314		28724	3.98	3.0E-25	IN 1282388	INT	Homo sapiens hypothetical protein FLJ20344 (FLJ20344), mRNA
3314	16074	28725	3.98	3.0E-25	TN 1233321 NT		Homo sapiens hypothetical protein FLJ20344 (FLJ20344), mRNA
4837	17568	30190	0.75			SWISSPROT	KALLISTATIN PRECURSOR (KALLIKREIN INHIBITOR) (PROTEASE INHIBITOR 4)
8518	10284	32288	90			HOT HIMAN	np27b02.s1 NCI_CGAP_Pr22 Homo sapiens cDNA clone IMAGE:1117515.3' similar to gb:M61866 ZINC FINCER PROTEIN 95 (HTIMAN)
8235	1	34065	4 86		3 0F-25 AI 163210 2	Т	Homo sariens chromosome 21 segment HS21(0)10
	1						m30h10.s1 NCI CGAP Pri Home serviens CDNA clane IMAGE-015331 similar to contains 1.1.11.1
10959	23635	36886	1.99	3.0E-25	3.0E-25 AA579013.1	EST HUMAN	repetitive element
1326	ŀ	26749	2.94	2.0E-25	2158		Homo saplens transducin (beta)-like 1 (TBL1) mRNA
2306	15031	27768	8.42	2.0E-25		EST_HUMAN	601511530F1 NIH_MGC_71 Homo sapiens cDNA clane IMAGE:3913087 5
2835		27897	3.67	2.0E-25		SWISSPROT	40S RIBOSOMAL PROTEIN S16
4167		29535	1.76	2.0E-25 P17008			40S RIBOSOMAL PROTEIN S16
4167	16907	29536	1.76	2.0E-25		SWISSPROT	40S RIBOSOMAL PROTEIN S16

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
1154	13909	26573	3.61		5.0E-26 AI708235.1	EST_HUMAN	as38h08.x1 Barstead aorta HPLRB6 Homo sapiens cDNA clone IMAGE:2319519 3' similar to WP:F49C12.11 CE03371;
1535	14282		1.4	4.0E-2	6 AA329548.1	EST_HUMAN	EST33448 Embryo, 12 week II Homo sapiens cDNA 5' end
9312	21979		3.72	4.0E-26	T857670 NT	N	Homo sapiens upstream binding transcription factor, RNA polymerase I (UBTF), mRNA
10558	23254	36491	2.75		4.0E-26 BE266187.1	EST_HUMAN	601191345F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3535210 67
1753	14495	27194			3.0E-26 D14547.1	F	Human DNA, SINE repetitive element
1896	14732	27464	1.31		3.0E-26 AL045855.2	EST_HUMAN	DKFZp4341066_11 434 (synonym: htes3) Homo saplens cDNA clone DKFZp4341066 51
2025	14760		3.15		3.0E-26 AA115895.1	EST HUMAN	2n30d08.rt Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens cDNA clone IMAGE:548943 5 similar to db:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN).
3760	16512	29148	1.04		3.0E-28 AA152464.1	EST_HUMAN	2030f10.r1 Stratagene colon (#937204) Homo saplens cDNA clone IMAGE:588427 5' similar to TR:G695374 G695374 THYROID RECEPTOR INTERACTOR:
3760	16512	29149	1.04		3.0E-26 AA152464.1	EST_HUMAN	2030/10.r1 Stratagene colon (#837204) Homo sapiens cDNA clone IMAGE:588427 5' similar to TR:G695374 G695374 THYROID RECEPTOR INTERACTOR:
6811	19472	32495	1.78		3.0E-26 BF245458.1	EST HUMAN	601864963F1 NIH MGC_57 Hamo saplens cDNA clone IMAGE:4083278 5'
10626	23319		1.42			Z	Homo sapiens MLL (MLL) gene, exons 1-3, and partial cds
11559	24158	37468	1.83		3.0E-26 AW875651.1	EST_HUMAN	QV2-PT0012-040400-124-e05 PT0012 Homo sapiens cDNA
11559	24158	37469	1.83		3.0E-26 AW875651.1	EST_HUMAN	QV2-PT0012-040400-124-e05 PT0012 Homo sapiens cDNA
11602	24201	37523	99'9		AA583173.1	EST_HUMAN	nn37d05.s1 NOI_CGAP_GC5 Homo saplens cDNA clone IMAGE:1086057 3' similar to contains OFR.t1 OFR repetitive element;
11858	24442	37783	1.36		3.0E-26 AF228925.1	NT	Mus musculus harmonin isoform b3 (Ush1c) mRNA, complete cds, alternatively spliced
12724	24995		2.52		3.0E-26 AW073434.1	EST HUMAN	xa57b09.x1 NC_CGAP_HSC2 Homo sapiens cDNA clone IMAGE.2570873 3' similar to contains MER30.t1 MER30 repetitive element;
999	13442	26083	92.9		2.0E-26 AL163282.2	NT	Homo sapiens chromosome 21 segment HS21C082
1861	14599		3.07			EST_HUMAN	DKFZp566L171_s1 566 (synonym: hfkd2) Homo saplens cDNA clone DKFZp566L171 3'
3225	15988	28642	5.89		2.0E-26 X86694.1	LΝ	M.musculus mRNA for astrocytic phosphoprotein, PEA-15
5147	17866		1.09			TN	Homo sapiens myotubularin related protein 7 mRNA, partial cds
10653	23344		2.7			NT	Homo sapiens DNA for amyloid precursor protein, complete cds
11180	23846	37132	6	2 OF-26		NAMIN TRA	to89a01.x1 NCL_CGAP_Gas4 Homo saplens cDNA clone IMAGE:2185416.3' similar to contains Alu
11395	24001		2.45	2.0E-26	T	LN	Homo saplens MHC class 1 region
12106	24603		1.57	2.0E-28	2.0E-26 AB037859.1	ĮŅ.	Homo sapiens mRNA for KIAA1438 protein, partial cds
133	12948	26691	5.18		Γ	EST HUMAN	QV4-HT0538-020300-123-602 HT0538 Homo sapiens cDNA
2040	14774	27503	1.37	1.0E-26	1.0E-26 AL039363.2	EST_HUMAN	DKFZp434H1910_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434H1910 5'
2693	15402		9:04	1.0E-26	П	LΝ	Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GADPH) mRNA, complete ods

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Top Hit Descriptor	602121491F1 NIH_MGC_56 Home sapiens cDNA clone IMAGE:4278527 5'	602121491F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4278527 5'	Mus musculus sparm tall associated protein (Stap), mRNA	Hamo sapiens chromosome 21 segment HS21C009	Rattus norvegicus putative four repeat ion channel mRNA, complete cds	QV0-OT0033-070300-152-b10 OT0033 Homo sapiens cDNA	H.sapiens DNA for endogenous retroviral like element	R.rettus RYA3 mRNA for a potential ligand-binding protein	PM0-BT0527-090100-001-d11 BT0527 Homo sapiens cDNA	7844C08 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7844C08	7e33f02x1 NCI_CGAP_Lu24 Homo saplens cDNA cione IMAGE:3284283 3'	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	Homo sapiens alpha NAC mRNA, complete cds	Inko1b10.s1 NCI_CGAP_Pr11 Homo sapiens cDNA clone IMAGE:1000699 similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 (HUMAN);	hi61h12x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2975879 3' similar to TR:076040 O76040 ORF2: FUNCTION UNKNOWN:	Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene	Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene	Rattus norvegicus voltage-gated sodium channel mRNA, complete cds	y39e01.1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:150840 6' similar to SP:HMGC_MOUSE Q02591 HOMEOBOX PROTEIN;	W28g07x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2426268 3'	nh08h05.s1 NCI_CGAP_Thy1 Home saplens cDNA clone IMAGE:943737 similar to contains L1.t3 L1 repetitive element:	R.rattus RYA3 mRNA for a potential ligand-binding protein	EST00738 Fetal brain, Stratagene (cat#636206) Homo sapiens cDNA clone HFBCF07	EST00738 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCF07	AU121685 MAMMA1 Homo sapiens cDNA done MAMMA1000746 5	nko1b10.s1 NCI_CGAP_Pr11 Homo sapiens cDNA clone IMAGE:1000699 similar to gb:M17886 60S ACIDIC RIBOSOMAL PROTEIN P1 (HUMAN);	11)1
Top Hit Database Source	EST_HUMAN	EST_HUMAN	F	NT	N	EST_HUMAN	LΝ	۲N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	EST_HUMAN	Z	Ł	¥	EST HUMAN	EST_HUMAN	EST HUMAN	N F	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN
Top Hit Acessian No.		BF666614.1	0569		AF078779.1	AW880859.1	X89211.1	X60658.1	BE071924.1	AA077705.1		BF035327.1	AF054187.1	AA565345.1	AW629172.1	AF111167.2	AF111167.2	AF000368.1	H02655.1	AI866347.1	AA551527.1		M78590.1	M78590.1	AU121685.1	AA565345.1	Ī
Most Similar (Top) Hit BLAST E Value	5.0E-27				4.0E-27	4.0E-27	4.0E-27	3.0E-27	3.0E-27		3.0E-27		2.0E-27		2.0E-27	2.0E-27	20E-27	2.0E-27				2.0E-27	2.0E-27	2.0E-27	2.0E-27	2.0E-27	
Expression Signal	2.82	2.82	1.65	1.07	1.54	0.7	1.98	4.61	1.06	6.24	0.63	2.93	14.84	5.12	10.39	1.96	1.96	1.36	0.61	1.65	23	0.76	1.28	1.28	4.11	3.31	1.57
ORF SEQ ID NO:			32421				37525	27499	29604	30697	33475					28632	28633	28385	32353	33810	,	35536	35782		36802		
Exan SEQ ID NO:	1 1	: 1	ļ	t I	20578	L	24203	14769		18068	П		12868	14625	15872	15981	15981	16755	19340		ı	22342	i	22583			1
Probe SEQ ID NO:	10137	10137	6645	7840	7883	9644	11604	2034	4238	5262	7698	9205	40	1888	3107	3218	3218	4009	6577	7989	9169	9891	9935	9935	10875	11469	428

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Top Hit Descriptor	Homo saplans DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	ht09g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3140256 3' similar to contains MER29.b3 MER29 repetitive element :	Homo saplens Retina-derived POU-domain factor-1 (RPF-1), mRNA	HSPD20461 HM3 Homo sapiens dDNA clone s4000095C10	HSPD20461 HM3 Homo sapiens aDNA clane 94000095C10	Homo saplens mRNA for KIAA0454 protein, partial cds	RC6-BT0627-140200-011-E06 BT0627 Harro saplens cDNA	Human mRNA for KIAA0260 gene, partial cds	Bos taurus letrophilin 3 splice variant bbah mRNA, complete cds	hw17c11.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3183188 3' similar to TR:Q07314 Q07314 SECRETED NEUREXIN III.AI PHA-C PRFCLIRSOR 131 TR-CA7243 1	AU126260 NT2RP1 Homo sapiens cDNA clone NT2RP1000443.5	zp18g12.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:60g8623	CM2-TN0140-070900-372-g01 TN0140 Homo saplens cDNA	au83h08.x1 Schnelder fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782911 3' almilar to TR:060302 O60302 KIAA0555 PROTEIN :conteins element MFR22 renefitive element :	AU142750 Y79AA1 Homo saplens cDNA clone Y79AA1000824 5'	Homo sapiens gamma-glutamytransferase-like activity 1 (GGTLA1), mRNA	AV735348 CB Homo saplens cDNA clone CBFAKA12 5'	Homo saplens zinc finger protein ZNF191 (ZNF191) gene, complete cds	ae60e03.r1 NCL_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:825340 6' elmilar to contains Alu repetitive element contains element PTR5 repetitive element:	wo18c07.x1 NCL_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2455692 3' similar to contains THR.b1	THR repetitive element;	y89f10.r1 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:146443 5'	xn33c09.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2695504 3' similar to SW:GG95_HUMAN Q08379 GOLGIN-95.:	Homo saplens myosin phosphatase, target subunit 1 (MVPT11), mBNA	601300703F1 NIH MGC 21 Homo sepiens cDNA clone IMAGE: 3635305 5'	qf86f10.x1 Scares_testls_NHT Homo sapiens cDNA done IMAGE:1755019 3' similar to gb:M19503 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (HUMAN);
Top Hit Database Source	Ϋ́	HEST HUMAN M		EST_HUMAN H	Г		EST_HUMAN R	Ĭ	NT B	hy EST HUMAN SE	Т	Г	Г	EST HUMAN TE	Г		EST_HUMAN AV		aa EST HUMAN reg	Г	EST_HUMAN TH	EST_HUMAN 1/8		Ĭ.	T HUMAN	HUMAN
Top Hit Acessian No.	1.0E-27 AB026898.1	1.0E-27 BE350127.1	5855	1.0E-27 F30158.1			1.0E-27 BE079780.1	Г			Γ			8.0E-28 AW157571.1	Γ	7866			AA504562.1		5.0E-28 AI921003.1		AW195066.1	5318	BE409100.1	Al198941.1 E
Most Similar (Top) Hit BLAST E Value	1.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27	1.0E-27	9.0E-28	9.0E-28	9.0E-28	9.0E-28	8.0E-28	7.0E-28	7.0E-28	7.0E-28	6.0E-28	6.0E-28		5.0E-28	6.0E-28			4.0E-28	
Expression Signal	1.34	0.98	6.26	1.96	1.98	0.98	2.28	2.65	3.51	2.94	3.31	0.63	4.85	2.48	7.89	3.36	2.78	0.97	2.35		4.19	1.44	1.68	0.78	2.52	1.88
ORF SEQ ID NO:	26404		32215	32542	32543	34346		36463	37625		25747	36150			26576	37089						28372	28087	28389	28511	32988
Exen SEQ ID NO:	13741	16806	19217	19515		1		22275	24299	12961	13107	22937	24504	25245	13913	23809	24474	21509	24873		13114	16738	15343	15742	15871	19915
Probe SEQ ID NO:	976	4061	6446	6771	6771	8208	8884	9622	11704	137	303	10289	11951	12268	1158	11142	11910	8817	12526		310	3880	283	2976	3106	7230

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
, in	13700		80 %	AC 30 A	A E020308 4	TN	Home satisfact chromosome 9 dunification of the T cell recentor beta locus and trocsingen name families
10928	1		17.24	4.0E-28	AB038241.1	IN	Felis catus GAPDH mRNA for glyceraldehyde-3-phosphate dehydrogenase, complete ods
10950	1	32988		4.0E-28	Al198941.1	EST HUMAN	qf86f10.x1 Soares, testis, NHT Homo sapiens cDNA clone IMAGE:1755019 3' similar to gb:M19503 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (HUMAN);
12312	24734			4.0E-28	AW854244.1	EST_HUMAN	RC3-CT0254-240400-210-f12 CT0254 Homo saplens cDNA
1260	l		2.88	3.0E-28	AF155382.1	TN	Homo sapiens metalloprotease-like, disintegrin-like, cysteine-rich protein 2 epsilon (ADAM22) mRNA, complete cds
5051	ı		1.05	3.0E-28	AF009660.1	NT	Homo saplens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region
8726	21418	34562	1.89	3.0E-28	BF354030.1	EST_HUMAN	MR3-HT0713-280500-013-f09 HT0713 Homo sapiens cDNA
10853	23533	36778		3.0E-28	U53588.1	LN	Homo sapiens MHC class 1 region
12344	24751		3.62	3.0E-28	AI831991.1	EST HUMAN	wj98f07.x1 NCI_CGAP_Lym12 Homo saplens cDNA clone IMAGE:2410885 3' similar to contains Alu repetitive element HGR repetitive element;
87	12913	25551		2.0E-28	BE062167.1	EST HUMAN	RC1-BT0254-220300-019-e05 BT0254 Homo sapiens cDNA
1023	13783		0.86		4501912 NT	LN	Homo saplens a disintegrin and metalloproteinase domain 23 (ADAM23) mRNA
1142	13897	26558	16.03	2.0E-28	Y11107.3	NT	Homo saplens ITGB4 gene for Integrin beta 4 subunit, exons 3-41
2481	15199		2.1	2.0E-28	Al348634.1	EST HUMAN	qos5b08.x1 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1910483 3' similar to contains L1.b2 L1 repetitive element;
							hr76c03 x1 NCI CGAP Kid11 Home sablens cDNA clone IMAGE:3134404 3' similar to contains LOR1.b1
6215	18989	31966		2.0E-28	BF224402.1	EST_HUMAN	LOR1 repetitive element;
6238			5.07	2.0E-28	BF212305.1	EST_HUMAN	601814196F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4048751 5'
7943	1	33766	0.71	2.0E-28	AF005273.1	NT	Sus scrofa domestica submaxillary apomucin mRNA, complete ods
9484 484	22137		5.54	2.0E-28	AW972305.1	EST_HUMAN	EST384394 MAGE reseguences, MAGL Homo sapiens cDNA
11814	24212	97536	184	90E-28	AF224669 1	LΝ	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (UBE2D3) genes, complete cds
12322	1			2.0E-28	H06376.1	EST_HUMAN	y/79c09.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:44300 5'
1461	14208			1.0E-28	D38044.1	NT	Human gene for Ah-receptor, exon 7-9
2217	14945	27685	2.37	1.0E-28	BF333236.1	EST_HUMAN	QV1-BT0821-120900-360-b03 BT0821 Homo sapiens cDNA
7759	20455		3.2		11429885 NT	LΝ	Homo sapiens similar to ribosomal protein L12 (H. sapiens) (LOC63091), mRNA
797	20612		3.3		8922793 NT	NT	Hamo sapiens hypothetical protein FLJ10968 (FLJ10968), mRNA
							EST179615 HCC cell line (matastasts to liver in mouse) II Homo sapiens cDNA 5' end similar to similar to
9178	21848	35014	4.84	1.0E-28	AA308744.1	EST_HUMAN	retroviral LTR
9776				1.0E-28		LN	Homo sapiens gamma-glutamytransferase-like activity 1 (GGTLA1), mRNA
9776	22427	35634	8.73		4758431 NT	L'A	Homo sapiens gamma-glutamyltransferasa-like activity 1 (GGTLA1), mRNA

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Top Hit Descriptor	Homo sapiens envelope protein RIC-6 (env) gene, complete cds	wr85d10.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2492563 3' similar to TR:015546 015546 HERV-E ENVELOPE GLYCOPROTEIN ;	wr65d10.x1 NO_CGAP_U11 Homo sapiens cDNA done IMAGE:2492563 3' similar to TR:O16546 O15546 HERVE ENVELOPE GLYCOPROTEIN :	Homo sapiens chromosome 21 segment HS21C068	os71e04.x1 NCI_CGAP_GC2 Homo sapiens cDNA clone IMAGE:1610814 3' similar to contains L1.t2 L1 repetitive element;	wt27g07.x1 Soares_NFL_T_GBC_S1 Homo saptens cDNA clone IMAGE:2356860 3' similar to contains element MER6 repetitive element;	wf27g07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2356860 3' shnilar to contains element MER8 repetitive element:	601442206F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3848648 5'	Homo sapiens DNA-binding protein (LOC56242), mRNA	Homo sapiens DNA-binding protein (LOC56242), mRNA	Homo sapiens chromosome 21 segment HS210048	Homo sapiens chromosome 21 segment HS21 CO48	Homo sapiens chromosome 21 segment HS210048	Homo saplens chromosome 21 segment HS21C048	601669934F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3952833 5'	Homo sapiens splicing factor similar to dnaJ (SPF31), mRNA	QV0-OT0032-080300-155-d01 OT0032 Homo saplens cDNA	RC1-HN0003-220300-021-b04 HN0003 Homo saplens cDNA	R.rattus RYA3 mRNA for a potential ligand-binding protein	nz20c07.s1 NCI_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:1288332 3' similer to contains MER4.b1 MER4 repetitive element :	Homo sapiens zincilron regulated transporter-like (ZIRTL), mRNA	HSC23F051 normalized infant brain cDNA Homo saplens cDNA clone c-23f05	EST97317 Thymus I Homo saplens cDNA 5' end similar to EST containing O family repeat	PT2.1_13_B11.r tumor2 Homo sapiens cDNA 3'	PM4-BT0724-150400-004-d11 BT0724 Homo saplens cDNA	Human mRNA for integrin alpha subunit, complete cds	QV0-BN0147-290400-214-f12 BN0147 Homo sapiens cDNA	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -
Top Hit Database Source	F	EST HUMAN	EST HUMAN	Π	EST HUMAN			Г			TN	L		IN	THUMAN		EST_HUMAN (EST_HUMAN F	NT TN	EST HUMAN		T_HUMAN	Г	EST_HUMAN F	T_HUMAN		EST_HUMAN (Г
Top Hit Acession No.	AF084869.1	A1963604.1	Al963604.1		AI082459.1				37821	10567821 NT	AL163248.2	AL163248.2			BF025947.1	5108		AW983880.1	X60658.1	AA761215.1	22745	F08688.1	AA383873.1	A1557072.1	BE091133.1	D25303.1		A E4777777 4
Most Similar (Top) Hit BLAST E Value	2.0E-29/		2.0E-29/		2.0E-29	_	_	-	2.0E-29	2.0E-29	2.0E-29/	2.0E-29	2.0E-29	2.0E-29 /	20E-29		2.0E-29/	1.0E-29 /	1.0E-29)	9.05-30			8.0E-30	8.0E-30	7.0E-30 E	6.0E-30	6.0E-30	4 0F-30 A
Expression Signal	1.72	6.62	8.62	1.63	0.99	1.48	1.36	1.16	0.63	9.63	3.61	3.61	3.61	3.61	1.31	2.04	1.73	7.37	0.85	2.97	1.76	8.94	3.72	3.1	1.03	1.73	2.3	0.48
ORF SEQ ID NO:	25901		26956	29610	31449	31830				34314	35241	35242	35999	36000	36804			34527	36391	32255			33994	34412			28598	l
Exen SEQ ID NO:	13265		14270	l	18527	18865	l		L.	21169			22787		23557		24100		23164	19254	24531	19001	20862	21275	14251			
Probe SEQ ID NO:	480	1523	1523	4246	6735	6087	7459	7878	8477	8477	9408	9408	10139	10139	10877	11459	11499	8691	10518	6487	11992	6227	8168	8283	1505	1766	3185	10437

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Single Exon Probes Expressed in Brain	Hit Asse Top Hit Descriptor	Human lambda-mmunoglobulin constant region complex (germline)		T	Homo saplens chromosome 21 section 14 2000 1	Homo saplens chromosome 21 segment HSZICMO	Homo sapiens chromosome 21 segment HS21C010	Г	Т	Г	Т		Т	Raftus norveolicus nursative four eneat inn channel militari sursative etc.	Rattus norvealcus buteitiva four repeat ion channel mayor, complete de		T	Homo saplens mRNA for KIAA1143 protein partial cytes		Г	Γ		Γ	Г	1	Г	Т			T
Single Exon	Top Hit Acession Database No. Source	1755.1 NT	30 Ai399992.1 EST HUMAN		Π		30 AL163210.2 NT	30 AW837471.1 EST HUMAN	Γ		-30 AW812488.1 EST HUMAN			30 AF078779.1 NT	30 AF078779.1 NT	30 BE350127.1 EST HUMAN			356 SWISSPROT	5.1		30 BE175877.1 EST_HUMAN	35232.1 EST_HUMAN	Γ	l		38945.1 EST_HUMAN			Ī
	Most Similar (Top) Hit BLAST E Value	3.38 6.0E-30 X51755.1	5.0E	5.0E	5.0E	5.0E	5.0E	4.0E	4.0E	4.0Ε	4.0E	3.0	3.0E-	3.0E	3.0E	3.0E	3.0E	3.0E-	3.0E	2.0E-	2.0E-								5 2.0E-30 AA019103.1	
	SEQ Expression O: Signal	3.	29376 26.19	Ϋ́	۲	37047 2.47		27599 1.72	27600 1.72	30560 0.63	34641 2.82	2.11	29128 0.93	0.58	0.48	36200 1.		36343 0.53		26077 0.9		26896 5.31			160 2.26			32435 0.55		
	Exan ORF SEQ ID NO:	17897	16742 28	25176	23485	23773 37		14869 27	14869 27		21495 34	13885		20547	21078	22980 362													21068 342	
	Probe SEC ID NO:	12769			10802		ľ	- (2139	6756		1129	3740			10333	Ш		\perp			_(_ [2920 1			_[- [8375 2	

Page 259 of 536 Table 4 Single Exon Probes Expressed in Brain

. Top Hit Descriptor	7637c12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284662 3' similar to SW:DHSA_HUMAN P31040 SUCCINATE DEHYDROGENASE [UBIQUINONE] FLAVOPROTEIN SUBUNIT PRECURSOR;	7837c12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284662 3' similar to SW:DHSA_HUMAN P31040 SUCCINATE DEHYDROGENASE [UBIQUINONE] FLAVOPROTEIN SUBUNIT PRECURSOR;	EST383657 MAGE resequences, MAGL Homo sapiens cDNA	ha33d08.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2875499 3' similar to contains THR.b3 THR repetitive element;	C18939 Human placenta cDNA (TFujiwara) Homo saplens cDNA clone GEN-570C01 5'	hd30b04;x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone iMAGE:2910991 3' similar to contains MER1.t3 MER1 MER1 repetitive element;	Homo sapiens chromosome 21 segment HS21 C003	ac77b08.s1 Strategene lung (#937210) Homo sapiens cDNA clone IMAGE:868599 3'	602022560F1 NCI_CGAP_Brn67 Homo sapiens cDNA done IMAGE:4157991 5'	EST186868 HCC cell line (matastasis to liver in mouse) II Homo sapiens cDNA 5' end	601809932F1 NIH_MGC_18 Homo saplens cDNA clone IMAGE:4040694 5'	CHR220532 Chromosome 22 exon Homo sapiens cDNA clone C22_728 5'	yc65e06.r1 Stratagene liver (#937224) Homo sapiens cDNA clone IMAGE:85570 5'	yc65e06.r1 Stratagene liver (#937224) Homo sapiens cDNA clone IMAGE:85570 5'	yf99b08.r.1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:30566 6' similar to gb:X12953 RAS- RELATED PROTEIN RAB-2 (HUMAN);	y/99b08.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE.30566 5' similar to gb:X12953 RAS-	RELATED PROTEIN RAB-2 (HUMAN);	HSC05F032 normalized Infant brain cDNA Homo sapiens cDNA cione c-05f03 3'	Rattus norvegicus putative four repeat ion channel mRNA, complete cds	Homo sapiens hypothetical protein FLJ20420 (FLJ20420), mRNA	Homo sapiens chromosome 21 segment HS21C008	OLFACTORY RECEPTOR 15 (OR3)	OLFACTORY RECEPTOR 15 (OR3)	hw05a11.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3182012 3'	hw05a11.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3182012 3'	Homo sapiens V1-vascular vasopressin receptor AVPR1A gene, promoter region and partial cds	Homo sapiens V1-vascular vasopressin receptor AVPR1A gene, promoter region and partial cds
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN	EST_HUMAN	FZ	۲	ΡZ	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	NT	۲
Top Hit Acession No.	2.0E-30 BE670617.1	BE670617.1	20E-30 AW971568.1	2.0E-30 AW470791.1	1.0E-30 C18939.1	1.0E-30 AW 468897.1	AL163203.2	AA664377.1	1.0E-30 BF347728.1	1.0E-30 AA315045.1	1.0E-30 BF183230.1	1.0E-30 H55593.1	9.0E-31 T73025.1	9.0E-31 T73025.1	9.0E-31 R18214.1		9.0E-31 R18214.1	9.0E-31 Z38293.1	9.0E-31 AF078779.1	TN 8923389 NT	8.0E-31 AL163208.2	P23275	P23275	7.0E-31 BE326617.1	7.0E-31 BE326517.1	7.0E-31 AF208541.1	
Most Similar (Top) Hit BLAST E Value	2.0E-30	2.0E-30	20E-30	2.0E-30	1.0E-30	1.0E-30	1.0E-30	1.0E-30	1.0E-30	1.0E-30				9.0E-31	9.0E-31		9.0E-31	9.0E-31	9.0E-31	8.0E-31	8.0E-31	8.0E-31 P23275	8.0E-31	7.0E-31	7.0E-31	7.0E-31	7.0E-31
Expression Signal	3.61	3.61	3.62	7.37	16.33	2.34	2.62	7.16	2.01	0.94	2.46	6.95	0.81	0.81	0.81		0.81	1.63	0.48	2.41	4.6	1.43	1.43	3.29	3.29	96.0	0.96
ORF SEQ ID NO:	34366	34367	35741	35839	25729	25942	26122	27675			33399		29135	29136	34053		34054		34359			30213	30214	28123		34130	34131
Exan SEQ ID NO:	21226	21225	L		13087	13309		L		15816	ı			16601	20917	1	20917	21214	21216	<u>L</u>	15135	17590	17590	ı	15383	ı	20994
Prabe SEQ ID NO:	8533	8533	2897	9982	280	525	88	5208	2464	3050	7624	12581	3748	3748	8223		8223	8522	8524	1054	2414	4861	4861	2674	2674	8300	8300

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Single Exon Probes Expressed in Brain Top Hit Describor	Single Exon Properties Source Source Source Source Source EST HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	Top Hit Acess No. No. 1 BE408611.1 X61765.1 AF055066.1 AF055066.1 BE30127.1 BE894486.1 M60694.1 M7073.1		Signal Signal 2.26 4.37 7.3 3.58 3.58 7.18 1.23 1.23 1.25 1.25 1.25 1.25 1.35 1.35 1.35 1.35 1.35 1.35 1.35 1.3	ORF SEQ ID NO: 26840 26841 26841 27040 27040 27056 27368 27368 27677 27800	Exon SEQ ID NO: 24805 24805 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 13000 14650 24872 20775 20775 20775 20776 14697 20776 20776 20776 14697 20776 20776 20776 20776 14697 20776 207776 207776 20776 20776 20776 20776 20776 20776 20776 20776 20776 20776 207776 20776 20776 20776 20776 20776 20776 20776 20776 20776 207776 207776 207776 20776 20776 20776 20776 20776 20776 20776 20776 20776 20776 207776 20776 20776 20776 20776 20776 20776 20776 20776 20776 20777776 207776 207776 207776 207776 2077776 207777777777	Probe SEQ ID NO: NO: 12434 12169 12169 12106 12206 12206 12206 12467 1239 10498 10498 10498 10498 10498 10498 10498 10498 10498 10527 11101 1101 1210 1210 1210 1210 1210 1
Control of the Control (Synanym: namyz) Home sapiens cDNA clone DKFZp761G1613 5'							
DKFZp761G1513_1 761 (synonym: hamy2) Home sapiens cDNA clone DKFZp7e1G1813_E	SI HUMAN		EVE 2	1	+	\mid	-
1971900.XI SORIES NFL 1 GBC S1 Home sapiens cDNA clone IMAGE:2111672 3/			20E 34	1 80	27800	15062	2339
MANAGE A SOCIETY AND THE SERVICE CONA	AT HIMAN		2.0E-31	1.09	27677	14939	2211
DV2. TORA SARAN 144 M3 1 TORA DE CONTROL DE	ST HUMAN	l	2.0E-31	1.37	27368	- <u>4</u>	DLA!
Homo sapiens mRNA for KIAA1342 protein partial cde	トラ		3.0E-31	8.		2000	2000
601458531F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3862088 R	ST_HUMAN		0.0E-01	20.5	+	25050	12810
40S KIBOSOMAL PROTEIN S15 (RIG PROTEIN)	OWISSPRO		2000	3 86		23771	11101
405 Pirocontain Seales Base NH Homo septens cDNA clone IMAGE:731047 6	TOGGGGGA	Ī	3.0F-31	2.04	36459	23224	10527
7 IORAGA 4 Society Letter 1 Transporter COS	NAMIN TS		3.0E-31	0.54	36371	23. 44.	10498
Horse mRNA for farith 1 - chain complete 1000	5		3.0E-31	3.7	35312	22132	947B
Homo sablens chromosome 21 segment US24.500.	F	7	3.0E-31	2		B	
Homo sapiens hypothetical protein FLJ10842 (FLJ10842), mRNA	Ę	8328 8328	3.05.31	310	+	20755	8
Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 8 (19kD, ASHI) (NDLFBR) mRNA		4440000	3 05 34	7	33151	20072	7393
More control of the state of th	Ę	4826853	3.0E-31	12.23	32999	18924	7239
From Septens KIAA0569 gene product (KIAA0569), mRNA							-
Troillo sapiens GG 1 gene, expn 1		02000	4 0F-34	186	F	24826	12467
Horse SETMAR) mariner transposase fusion gene (SETMAR) mRNA	Ė	AJ230125 1	4.0E-31	1.86		24672	12205
Home saviane CET down of a grant 102 10000	LN		4.0E-31	1.23		10,00	78/7
Homo saplens chromosome 21 segment HS34C000	Z	AL163280.2	4.0E-31	1.57		3	
ACETYLGALACTOSAMINYLTRANSFERASE) (GALNAC-71)	SWISSPROT		4.0E-31	E.O.		14550	1840
POLITIEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE) (I IDD. CALINA COLLARITIES IN					9	6	800
Homo sapiens Xq pseudoautosomal region; segment 1/2	2	T			-		
Control of Poco Elemen 1. Contains L1.t1 L1 repetitive element;	LIVANOI - LIV		4 OF-34	5.18		13362	582
7k06f04x1 NCI_CGAP_GC6 Homo septens aDNA clone IMAGE:3443479 3' similar to TR:Q13537 Q13.	EST HIMAN	BF056540.1	6.0E-31	0.73		21037	8344
Homo seplens type I DNA topolsomerase gene. exon 8	ĻΝ	M60694.1	3.UE-31	3.38	1		
Homo sapiens type i DNA topoisomerase gene, exon 8	2	1,0000	200	2000	ı	13000	187
T	TO LONG	Meneou 1	5 0E 34	3.58		13000	187
T	EST DIMAN	BER9448R 4	6.0E-31	1.96			12169
	EST HUMAN	BE350127.1	. 6.0E-31	99.0			8229
Homo saplens MHC class 1 region	TN	1 AF055066.1	6.0E-3	4.37		_	3
spliced	ΙN	1 AF223391.1		2.00		L	8063
Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cris alternation	1	A E222204 4		89 6			3667
	LN LN	1 X51765.1	7.0	2.26	1	- 1	12434
	EST_HUMAN	1 BE408611.1		18.0°	l	J	1
T		1 BEADBOAR A		0.94			9168
	Source		SCASTE Value		j j		ë
							SEQ ID
robes Expressed in Brain	ngle Exon P	ະ ເ ັດ					

PCT/US01/00667

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WO 01/57275

		Γ	3	12	Τ	Τ	Τ		Γ	Γ			T	Ţ		Γ	Τ	Τ	Τ	Τ	<u> </u>	T	Τ	7		_	T	Т	П
Single Exon Probes Expressed in Brain	Top Hit Descriptor	UI-H-Bi3-akb-f-09-0-UI.s1 NCI_CGAP_Sub5 Hamo sapiens cDNA clone IMAGE:2733833 3'	ht09g01.x1 NCI_CGAP_Kid13 Homo sepiens cDNA clone IMAGE:3146256 3' sImilar to contains MER29.b3 MER29 repetitive element;	nr06f04.s1 NCI_CGAP_Co10 Hamo saplens cDNA clone IMAGE:1161065 3' similar to TR:Q13537 Q13537 MER37 TRANSPOSABLE ELEMENT, COMPLETE CONSENSUS SEQUENCE:	Homo sapiens B9 protein (B9), mRNA	AV710948 Cu Hamo sapiens cDNA clane CuAALB07 5	AV710948 Cu Homo sapiens cDNA clone CuAALB07 5'	601304125F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638310 5'	601304125F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638310 5'	Homo sapiens hexakinase il gene, promoter region	HA1110 Human fetal liver cONA library Homo sapiens cONA	Homo sapiens MAGE-B2 (MAGE-B2), MAGE-B3 (MAGE-B3), MAGE-B4 (MAGE-B4), and MAGE-B1	(WAGE-51) genes, complete cds	OF ACCOUNT MEDICAL CONTROL OF ACCOUNT MEDICAL CO	OLFACTORY RECEPTOR 2C1	OLFACTORY RECEPTOR 2C1	DKFZp547B235_r1 647 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547B235 5'	DKFZp547B235_r1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp547B235 5'	MR3-ST0220-151299-028-a08_1 ST0220 Homo sepiens cDNA	Homo saplens minisatellite ceb1 repeat region	Bos taurus xenobiotic/medium-chain fatty acid:CoA ligase form XL-III mRNA, nuclear mRNA encoding imitochondrial protein, complete cds	Homo sapiens MAGE-B2 (MAGE-B2), MAGE-B3 (MAGE-B3), MAGE-B4 (MAGE-B4), and MAGE-B1 (MAGE-B1) ages, complete cds	qr21h03.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:1750709 3' similar to TR:Q16595	Q16595 FRATAXIN.;	Human germline T-cell receptor beta chain TCRBV17S1A1T, TCRBV2S1, TCRBV10S1P, TCRBV29S1P,	[TCRBV19S1P, TCRBV16S1, TCRBV11S1A1T, HVB relic, TCRBV28S1P, TCRBV34S1, TCRBV14S1, TCRBV34S1, TCRBV4S1, TCRBV3S1, TCRBV4S1, TCRBV3S1, TCRBV4S1, TCRBV3S1, TC	AV723976 HTB Homo sepiens cDNA clone HTBAAG01 5	Homo sapiens hypothetical protein FLJ11294 (FLJ11294), mRNA	oz15a09.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1675384 31
gie Exon Pro	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN FN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	į	NI	TOURS INC.	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	LN			EST HUMAN		F	EST HUMAN	NT	EST_HUMAN
	Top Hit Acession No.	2.0E-31 AW444498.1	2.0E-31 BE350127.1	31 AA877764.1	7661535 NT	31 AV710948.1	31 AV710948.1		31 BE408611.1		2.0E-31 AI114527.1		31 093163.1	1360	095371	095371	1.0E-31 AL134376.1	31 AL134376.1	1.0E-31 AW391679.1		1.0E-31 AF126145.1			31 AI086434.1		1 OE-34 188081 4		3082	8.0E-32 A1056770.1
	Most Similar (Top) Hit BLAST E Value	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	2.0E-31	1 07 04	1.05.4	2 1	1.0E-31 095371	1.0E-31	1.0E-31	1.0E-31	1.0E-31	1.0E-31	1.0E-31	1.0E-31		1.0E-31		1 OF.34	9.0E-32	9.0E-32	8.0E-32
	Expression Signal	0.81	3.57	2.05	3.64	0.94	0.94	2.35	2.35	1.58	1.75	77	11.09	31	1.35	1.35	1.15	1.15	3.79	2.2	-	0.51		2.7		1 48	2.38	99.0	2.48
	ORF SEQ ID NO:	30624	31334		34959	35661	35662	35829	35830			00710	20400	2002	27093	27094	28952	28953	30640	31782	32948	35995		36757		37752	32312		27530
	Exan SEQ ID NO:	18001	18421	21685	21795	22457	22457	22623	22623	24632	25413	10040	14404	5	14404	14404	17327	17327	18018	18822	19875	22784		23515		24414	18307	20164	14802
	Probe SEQ ID NO:	5193	5624	8975	9107	9086	9806	8975	8975	12144	12279	4	185	3 5	1638	1658	4592	4592	5210	6042	7189	10136	1	10833		11830	6542	7492	2070

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Exon ORF SEQ Expression Top Hit Acession Top Hit No. Signal BLAST E No. Delabase Source No. Signal BLAST E No. Delabase Source No. Signal Signal Source Signal Signal Source Sourc							_							_															
Exon ORF SEQ Expression CTOP) Hit Top Hit Accession Top Hit Top Hit Accession Delabase Signal BLASTE No. Signal BLASTE No. Signal BLASTE No. Signal BLASTE No. Source Value Signal Source Value Signal Source Value Signal Source Signal Source Sou	bes Expressed in Brain	Top Hit Descriptor	RC2-BN0048-201300-045-04 BN0048 U	NUCLEAR ENVELOPE PORE MEMBRANE PROTEIN POM 121 (PORE MEMBRANE PROTEIN OF 121) (P145)	Human chromosome 22 Immunoglobulin V(K)I gene, part. with 5' breakpoint between orphon and neighbouring non-smallful contracts.	bm34410.xt NCI_CGAP_Kid11 Home saplens cDNA clone IMAGE:2169994 3' similar to contains MER29.t3	AND EASTERNIES BETTENING TO THE STATE OF THE	Home college BB04464 BB1464	Homo septens if NOTIFIT mRNA, complete cds Homo septens chromosome 21 scannost Douglass	ws08h12.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2496847 3' similar to centains MER18.b3	WENTS Repeatable element;	nomo sapiens A i -binding transcription factor 1 (ATBF1), mRNA	Homo sapiens AT-binding transcription factor 1 (ATBF1), mRNA	KC4-B10311-141199-011-h06 BT0311 Homo sapiens cDNA	Homo sapiens FLI-1 gene, partial	AV 21000 HIP Homo sapiens cDNA clone HTFAKC07 5'	AV Jacobs Bin Homo saplens cDNA clone BMFBBH12 5'	AV 7880A BM Homo sapiens cDNA clone BMFBBH12 5' 295a07.st Soeres_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:448500 3' similar to	contains I HR.t3 THR repetitive element;	or recessor i vin Juleo 21 Homo sapiens oDNA clone IMAGE:3139701 6' Homo sapiens myeloid/iympholid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4 Mil 73 meNa	demosymments mysioid/lymphold or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to. 4	MILL 14) MINNA MATERIAL MIN 1000 CATE	Image cell 40 lines and a line septemb cDNA clone IMAGE:3139701 6	Septens mRNA for missing	Septembrilly for mystering	HBBANB of Stepanson Lie and a software in	n88008 r1 Stratagne Hells cell so 937216 Homo caplens cDNA clone IMAGE:563150 5	12056t Testis 1 Homo serions CDNA class 147056	AV736499 CB Homo septents cDNA clone CBFBIA08 5'
Exon DRF SEQ Expression Top) Hit Top Hit Acess No. Value No. No. Signal DRO: Signal DRAST E No. Value No. Value No. No	igle Exon Pro		EST HUMAN	SWISSPROT	L	MANI III TOB	EST HIMANI	LN FIN	LN	TOT TOU	TI TOWAN		ı			Т	T	T^-	1	NUMBER OF THE PROPERTY OF THE		E INVANI	Т	7-		H HUMAN	Т	Т	П
SEQ NO ORF SEQ Signal Nost Sin	יאַ ו	Top Hit Acession No.	AW897214.1	P52591	X17283.1	AI478104 1	BE88018 1	AF116827 1	AL163248.2	A IORFFOR 4	11430574	44400674	270/4			T	T		T	74574	5174574					Ļ	Γ	Γ	\prod
SEQ ID ORF SEQ Express			8.0E-32	7.0E-32	7.0E-32	6.0E-32	6.0E-32	5.0E-32	4.0E-32	4.0F-32	4.0E-32	4 OF 32	4.0E.32	20 TO C	3.0E-32	3.0F-32	3 0F-32 /	0 10 6	3.0E-32	3.0E-32	3.0E-32	3.0E-32 B	2.0E-32 N	20E-32Z	20E-32 Z	2.0E-32 A	20E-32 A	2.0E-32 T	2.0E-32 A
SEQ ID ORI		Expression Signal	0.97	0.98	8.19	1.01	1.47	16.78	1.78	0.91	2.94	284	12	100	8.08	8.38	8.38	2 57	3.51	2.97	2.97	2.27	0.81	5.32	5.32	2.28	2.28	2.96	2.42
8		ORF SEQ ID NO:				28180		26431			33266	33267		25872	28870	35135	35136	36768		30586	30587		31902	32155	32156	34003	34004	37784	30961
SEG ID NO: NO: 5384 4807 12122 12128 16148 16158 16168			LI				19950	13771	13877	17867	20174	20174	20951	13229	14 184	21961	21861	23528	24834	17899	17899	24958	18935	19158	19158	20870	20870	24443	26022
		Probe SEQ ID NO:	5394	4807	12122	2735	7266	1011	910	5148	7503	7503	8257	443	1437	9294	9294	10843	12148	12507	12507	12656	6158	6387	8387	8176	8176	11859	12763

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Top Hit Descriptor	AV736449 CB Homo saplens cDNA clone CBFBIA08 6'	601573207F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3834433 5'	Hamo sapiens chromosome 11open reading frame 9 (C11ORF9), mRNA	nw21g02.s1 NCI_CGAP_GCB0 Homo sapiens cDNA clone IMAGE:1241138 3' similar to contains THR.t3	THR repetitive element :	hw07c05.x1 NCI_CGAP_Lu24 Homo septens cDNA clone IMAGE:3182216 3' similar to TR:O88539 O88539 WW DOMAIN BINDING PROTEIN 11.;	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	602021164F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4156670 5	Homo sapiens chromosome 21 segment HS21 C080	Homo sapiens short-chain alcohol dehydrogenase family member (HEP27) mRNA	Homo sapiens short-chain alcohol dehydrogenase family member (HEP27) mRNA	to12b09.x1 NCI_CGAP_U2 Homo sapiens cDNA clone IMAGE:2178809 3' similar to contains OFR.t1 OFR	repetitive element;	AV730056 HTF Hamo sapiens cDNA clane HTFAVE08 5'	EST383396 MAGE resequences, MAGL Homo sapiens cDNA	Human HLRP mRNA for leukocyte common antigen-related peptide (protein-tyrosine phosphate) (EC 3.1.3.48)	602021164F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMACE:4156670 5	EST383657 MAGE resequences, MAGL Homo sapiens cDNA	no16h01.s1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100881 3' similar to contains L1.t1 L1	repetitive element ;	Horno sapiens chromosome 21 segment HS21C085	HSPD21201 HM3 Homo saplens cDNA clone s4000107H06	HSPD21201 HM3 Homo saplens cDNA clone s4000107H06	Human glyceraldehyde-3-phosphata dehydrogenase (GAPDH) gene, complete cds	Homo sapiens similar to RAD23 (S. cerevislae) homolog B (H. sapiens) (LOC63277), mRNA	Mus musculus SRY-box containing gene 6 (Sox6), mRNA	Mus musculus SRY-box containing gene 6 (Sox6), mRNA	QV1-FT0169-100700-271-a02 FT0169 Homo sapiens cDNA	Homo sapiens solute carrier family 5 (choline transporter), member 7 (SLC5A7), mRNA	Homo saplens spermidine synthase (SRM) mRNA	Homo sapiens spermidine synthase (SRM) mRNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	F		EST_HUMAN	EST_HUMAN	Į	EST HUMAN	Z	Z	E		EST_HUMAN	EST_HUMAN	EST_HUMAN	F	EST HUMAN	EST HUMAN		EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	FZ	N	Ł	Į,	EST_HUMAN	TN	۲N	LN
Top Hit Acession No.	2.0E-32 AV736449.1	BE743299.1	11439789 NT		1.0E-32 AA720574.1	9.0E-33 BE327112.1	9.0E-33 AF223391.1	9.0E-33 BF347229.1	9.0E-33 AL163280.2	5031736 NT	5031736 NT		7.0E-33 AI590115.1	7.0E-33 AV730056.1	7.0E-33 AW971307.1	7.0E-33 X54890.1	BF347229.1	7.0E-33 AW971568.1				6.0E-33 F30631.1	6.0E-33 F30631.1	J04038.1	11429198 NT	TN 60555609 NT	TN 6095249	5.0E-33 BF373515.1	11141884 NT	4507208(NT	4507208 NT
Most Similar (Top) Hit BLAST E Value	2.0E-32	1.0E-32	1.0E-32		1.0E-32	9.0E-33	9.0E-33	9.0E-33	9.0E-33	7.0E-33	7.0E-33		7.0E-33	7.0E-33	7.0E-33	7.0E-33	7.0E-33	7.0E-33		7.0E-33				6.0E-33 J04038.	6.0E-33	6.0E-33	6.0E-33	5.0E-33	5.0E-33	5.0E-33	5.0E-33
Expression Signal	2.42	1.67	7.02		8.08	4.68	4 05	1.95	5.22	3.14	3.14		2.29	6.45	9.3	1.56	2.41	1.93		4.34	0.94	96'0	96.0	9.33	3.09	1.12	1.12	1.46	1.19	1.43	1.43
ORF SEQ ID NO:	30962		32453		34329			34523		25517	25518		27622					37162		31090		31720	31721	34315	34438	35754	35755				27338
Exon SEQ ID NO:	25022				21186	16230	19096		23392	ı	12887		14888	ı		21537	1				16473	•	l			!	ı	1	Ι.	14628	L
Probe SEQ ID NO:	12763	3090	6955		8494	3474	6328	8687	10701	58	83		2158	2655	3236	8845	10732	11213		12127	3720	5976	5976	8478	8603	9910	9910	1770	1874	1891	1891

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	Top Hit Descriptor	Homo sapiens X-linked anhidroitic ectodermal dysplasia protein gene (EDA), expriz and iraniang report	regions	Homo sapiens calcium/calmodulin-dependent protein whase in Commercial	Homo sapiens F-box protein FBL4 (FBL4) mKNA, complete cus	Home sapiens protein kinase C beta-li type (PKKCB1) mixiva, curiptede cus	Human dystrophin (DMD) gene, exons 7, 8 and 9, and partial cds	OV3-BN0047-230200-102-b03 BN0047 Homo sapiens cDNA	Hirman dystrophin (DMD) gene, exons 7, 8 and 9, end pertial cds	westerne vi NCI CGAP Kid11 Homo sapiens cDNA clone IMAGE:2462410 31	Home sanians X-linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	AV727809 HTC Homo sapiens dury cigner in concile	Homo sapiens Xq pseudoautosomal region; segment 1/2	Homo seplens hypothetical protein FLJ10900 (FLJ10900), mRNA	MR4-BT0399-200100-001-h03 BT0399 Homo sapiens cDNA	art Farts it Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:108320 5	NATIONAL Spares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:108320 5	V14c10.11 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:148722 5	Human G2 protein mRNA, partial cds	Hirman G2 protein mRNA, partial cds	Museulus DAB/2J hair-specific (hacl-1) gene	Home saplens Nav38-binding protein Nav8P (LOC51729), mRNA	Human splicing factor SRp55-1 (SRp-55) mRNA, complete cds	Raftus norvegicus putative four repeat Ion channel mRNA, complete cds	Homo sapiens mRNA for KIAA1435 protein, partial cds	Homo sariens chromosome 21 segment HS21C009	HALAGE VAING CGAP Pr28 Homo sapiens cDNA clone IMAGE:2249194 3'	JOSEPH ST. Spares festis NHT Homo saplens cDNA clone IMAGE:1407936 31,	and 2000 HTM MGC 54 Homo saplens cDNA clone IMAGE:4102213 6	OUT TO THE TANK TO THE PROPERTY OF THE TOTAL TO THE TOTAL THE TOTA	Numer by some more and an experience of the control	601436331F1 NIT MOO CONTRIBUTION OF THE CONTRI	Wasguost Scarce in E. Control MER29.12 MER29.12 MER29 repetitive element;	
dy sogoi i iny alimin	Top Hit Database Source		IN					LI INVANI	Т	146741	EG TOMAN		EST HUMAN	Z L	TIN	EST HIMAN	TOT TOWNS	TOT TOT	EST HIMAN	FIGURE PLAN	111	2 2	Z	2 2	Į.	Z 12	i i		TOT HOMAN	EST HOMEN	ESI HOMAIN	L	EST_HUMAN	EST_HUMAN	
Z IIC	Top Hit Acession No.		-33 AF003528.1	4502556 NT	-33 AF199420.1	142075 4	A1397 3.1	J60022.1	-33 AW990610.1	-33 U60822.1	1.0E-33 AI927191.1	1.0E-33 AF003528.1	4 OE 33 AV727809 1	0 0E.34 A.1271735.1	TIM POCCOO	892273	8.0E-34 BE069882.1	=-34 T70845.1	E-34 T70845.1	7.0E-34 H12866.1	6.0E-34 U10991.1	6.0E-34 U10991.1	U03686.1	TIN DOCOT	5.0E-34 U30003.1	5.0E-34 AFU/8/79.1	5.0E-34 AB037856.1	AL163209.2	E-34 A1804667.1	E-34 AA861773.1	E-34 BF209778.1	E-34 M37277.1	E-34 BF035327.1	E-34 AI678101.1	
	Most Similar (Top) Hit BLAST E Value		1.0E-33/	1 OF 33	1 OF 33 /	4 00 00 4449075 4	1.05-50	1.0E-33 U60822.1	1.05-33/	1.0E-33	1.0E-33	1.0E-33	4 05 33	0 OF 34	9.00	8.0E-34	8.0E-34	7.0E-34	7.0E-34	7.0E-34	6.0E-34				5.0E-34				4.0	4.0	4.0	3.0	3.0	2.0	
	Expression Signal		7	2.48	840	0,0	1.04	0.84	1.83	3.32	2.21	407		1.41	1.0	0.98	0.66	2.27	0.56	3.05			2.13	2.9				1.79	1.64		1.28	0.78	3.14	1.16	
	ORF SEQ ID NO:	+		30,00	30,400	31180	33067		37251	37581				31005				28858	26858			25885	31107				38482	ic.	7 27449	1 31488	34769	1		1 34687	
	Exon SEQ ID NO:			5537	1/8/3	18299	19990	25432	23953	24259	L_	70007	ı	ł	25034	L	20353	L	<u> </u>	24662	13243	13243	L.	14611	17725	21457	23246	1_	1_	18561	L	ı			١
	Probe SEQ ID NO:		,	2	5158	5501	7307	9920	11292	11663	12407		125/0	12602	12780	2168	7689	1426	0066	12191	458	458	12011	1873	2005	8765	10550	11219	<u>8</u>	5770	8038	8438	11100	8850	;

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	wd35g06.x1 Soeres_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2330170 3' sImiler to contains	MICHARACTURACE PERGUNA Element	PLEXINA TRECURSOR (TRANSMEMBRANE PROTEIN SEX)	ADP ATP CARRIER PROTEIN, LIVERISOFORM T2 (ADP/ATP TRANSLOCASE 3) (ADENINE	Homo sepiens X-linked anhidrolite eotodermal dysplasia protein gene (EDA), exon 2 and flanking reneat	Successions	Home sapiens W.N.13 precursor (WNT3) mRNA, complete cds	POR PTOROS ANAMA SALAN S	NOZ-5 I USUG-Z4U4UU-U16-h08 BT0506 Homo sapiens cDNA	SOLINGTANDE VIII. VICE 69 Home seplens cDNA clone IMAGE:3888899 5	OLEACEONY FIGURE 69 Homo saplens cDNA clone IMAGE:3886999 5	OCTACIONY RECEPTOR-LIKE PROTEIN F5	ON 12500441003_11 584 (synonym: hfbr2) Homo saplens cDNA clone DKFZp584A1683 g/	0014/0382F1 NIH_MGC_67 Home sapiens cDNA clone IMAGE:3873476 5'	OU 14/0382F1 NIH _MGC_67 Homo saplens cDNA clone IMAGE:3873478 6	oc31c11.s1 NCI OGAP GCB1 Homo seniers CNN A CASTACT ASSESSED	TYROSINE-PROTEIN KINASE RECEPTOR FITA PRECIDE OF ALLIAMA.	Homo saplens chromosome 21 segment HS21C010	hh77b08.y1 NCI_CGAP_GU1 Homo saplens cDNA chose IAACE-30682537 FI	Homo saplens prohibitin (PHB) mRNA	nea33a08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3258134 3' similar to TR:O76912 O75912 DIACYLGLYCEROL KINASE IOTA	nae33a08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3258134 3' similar to TR:O75912	ONSWIZ DIACYLELYCEROL KINASE IOTA.;	PO TOWSCOOT I NIH, MGC_18 Home sapiens cDNA clane IMAGE:4040324 5	001230480F1 NIH_MGC_44 Homo saplens cDNA clane IMAGE:3608513 5	Homo English - L. C. C. Homo saplens cDNA clone IMAGE:4300860 3'	ahsahna at Seess 1-11 Till Till Till Till Till Till Till Ti	Homo septems vinc fineer models 200 / 2017 2017 200 / 2	UI-H-BW0-gld-4-09-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2731433 3'
gle Exon Prot	Top Hit Database Source	HST HIMAN	Т	Т		П			HIMAN	Т	Т	T	Т	7	EST HUMAN	NAME -		T_HUMAN		T_HUMAN		EST_HUMAN O		7	7	7	NONCL	T H! IMAN		T HUMAN
Sin	Top Hit Acession No.	E-34 Al678101.1					1.0E-34 AY009397 1		T	T	Ī		5.0	T	T	30500		1.0E-34 AA807097.1 E		9.0E-35 AW 663302.1 E	6031190 NT	-35 BF589937.1 E	-35 BF589937 4 E	T	1	T	5417	35 AA757115.1 F	5975	35 AW297191.1 ES
Most Silver	Most Similar (Top) Hit BLAST E Value	2.0E-34	2.0E-34 P51805	2.0E-34 P51805	1.0E-34 P12238	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34	1.0E-34 P23268	1.0E-34	1 0F-34 F	1.0E-34 P	1.0E-34		1.0E-34 A	1.0E-34 A	9.0E-35 A	8.0E-35	8.0E-35 B	8.0E-35 Br	8.0E-35 Br	8.0E-35 BF	8 OF 35 B	7.0E-35	6.0E-35 A	6.0E-35	8.0E-35 AV
	Expression Signal	1.16	1.34	1.34	6,53	1,32	76.0	76.0	3.44	2.05	2.05	0.45	7.1	1.39	1.39	1.82	-	1.65	422	7.	7.74	3.43	3.43	2.69	2.42	3.85	2.05	1.83	2.09	0.84
	ORF SEQ ID NO:			37058	26928	29055	29425	29426		31788	31789	35076	35434	37083	37084	37100	-	+	00000	20028	+	27171	27172	30170	36522	-	32163	26815	27409	29406
	SEQ ID NO:	1	_!	23783	14241	18416	16796	18796	17182	18827	18827	21904	22249	23805	23805	23820	02020	2000	18380	2000	8700	14472	14472	17545	23283	24611	19162	14138	14696	16775
	SEQ ID NO:	8850	11113	11113	1494	3663	4051	4051	4446	6047	6047	9225	9296	11138	11138	11153	12272	12502	3636	248		1730	1730	4814	10589	12119	6383	1391	1960	4030

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	Top Hit Descriptor	(TRIO), mRNA	nomo squene upo incomercia.	H.saplens mink I all order from the control of the	H. saplens mich and a contraction of the contractio	Human mRNA for KIAA0366 gene, partial cds	Homo saplens mRNA for KIAA1365 protein, partua cas	H.sapiens Immunoglobulin kappe light chain vartable region L.14	Homo sapiens mRNA for KIAA0406 protein, partial cds	Homo sapiens Ring1 and YY1 binding protein (RYBP), mRNA	Home sablens cli/2 kinase (CLX2), propin1, cote1, glucocerebrosidase (GBA), and metavin genes, complete Home sablens cli/2 kinase (CLX2), propin1, cote1, glucocerebrosidase (GBA), and metavin genes, complete the sablens cli/2 kinase (GBA).	cds; metazin pseudogene and glucocarebrosidase pseudogene; and thrombospondins (Tribos) years, paramic	cds	001431904F1 Nun_woo	9938605.X1 Sogres Jesus, virgin 1977.9 PROTEIN KIAA0249. ; SW:Y249 HUMAN 092539 HYPOTHETICAL PROTEIN KIAA0249. ;	lad38c05.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1837448 3 similar to	SW:Y249_HUMAN Q92539 HYPOTHETICAL PROTEIN KIAA0249:	ANSTALL Societies Tekin lives special in the land lives lives the land lives lives the land lives lives the land lives lives the land lives lives the land lives lives the land lives the	601109719F1 NIH_MGC_16 Homo Sapiens curs curio intro-contains	N198e07.r1 Sogres fetal liver spleen 1NFLS Homo sapiens CDNA clone INNACE.241230 Comment	PTR5 repetitive element; However amiens X-linked anhidrotitic ectodernal dysplasta protein gene (EDA), exon 2 and flanking repeat	regions regions C:442 Homo serviens CDNA clone IMAGE:3146256 3' similar to contains MER29.b3	httogori XI Not_Cook_ Nation to the second of the second of the File of the Fi	DKFZp434L148_r1 434 (synonym: https:// Homo eaplens curve curve Dry types 11 434 (synonym: https://	XV17703.X1 Soares_NFL_T_GBC_S1 Hamo sapiens cDNA clone IMAGE.2813405 3 SIMILE W CONTROLLY	repetitive element;contains MER19.r2 MER19 repetitive element;	9011222011 INITIATION From hare 1 dene complete cds	Home september of the TR. 1992 April 1994 Home september CDNA clone IMAGE:35653613's similar to TR. 1992 APRIL 1994 Home september CDNA clone IMAGE:35653613's similar to TR. 1992 APRIL 1994 April 19	1902DET - BOX PROTEIN FBI.2. : CONTROL OF STATE	Q9QZH7 F-BOX PROTEIN FBL2.;
an i liova Bigino	Top Hit Database Source				N	NT.	IN.	±N	FZ		-			EST_HUMAN	EST HIMAN		EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN	Ŗ	FST HUMAN	EST HIMAN			EST HUMAN	Ł	EST_HUMAN	EST_HUMAN
BIIIO	Top Hit Acession No.		6005921 NT	X94232.1	X94232.1	4.1	Ī	T	,	2020	202180		5.0E-35 AF023268.1	5.0E-35 BE890992.1	1 2000	5.0E-35 AIZU8/05.1	5.0E-35 AI208765.1	AA001786.1	4 of 36 BE257007 4		4.0E-35 H91193.1	15 AF003528.1	ac DE350427 1	A1 0 4650 A	35 ALU40390.1	35 AW303317.1	BE268182.1	3.0E-35 AF224492.1	-35 BF433100.1	-35 BF433100.1
}	B + m	value	6.0E-35	6.0E-35 X				0.05-30-0	5.05-30	5.0E-30.A	5.0E-35		5.0E-35	5.0E-35	100	5.0E-30/	5.0E-35	5 0F-35	30 30 7	4.0E-30	4.0E-35	4.0E-35	30 10 7	4.05-0	4.0E-35	4.0E	3.0E		3.0E	3.0E
	Expression Signal		3.84	0.93	000	26.0	0.00	3.17	1.36	1.07	1.7		1.7	3.99		2.35	235	37.0	24.4	16.80	4.87	0.72		T8.T	8.88	1.38		1.5	31.47	31.47
!	ORF SEQ ID NO:	_	33615	24445	97776	34440	35403	35656	27146	28232	28424		29746	1		33936	20007			26845	27265				34248	37646			30690	
	SEQ ID	į	20,00	2000	71307	21302	22218	22454	14447	15492	15775		17113	L		20803	i	- 1		14161	14551		L	19796	3 21109	24322	L		L_	
		į	1100	08/	200	8610	9565	9803	1704	2787	3008		4378	2 2	500	8109		8109	11130	1413	1811	7,760	Š	7108	8418	77.30	4570	2330	8258	5256

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Table 4

Proble (EAD) EAD (SEC) Characteristic (Top) His Accession (Top) His						<u>ਨ</u>	igle Exon Pro	Single Exon Probes Expressed in Brain
8 22050 1.42 3.0E-35 AP223391.1 NT 4 22722 35639 1.12 3.0E-35 AW003063.1 EST_HUMAN 5 16535 2.5667 1.88 2.0E-35 N88965.1 EST_HUMAN 1 16540 2.5687 1.65 2.0E-35 AB018413.1 NT 1 16066 2.8714 1.12 2.0E-35 AB020702.1 NT 1 16066 2.8715 1.12 2.0E-35 AB020702.1 NT 1 1630 2.8714 1.12 2.0E-35 AB020702.1 NT 1 16640 2.82280 0.78 2.0E-35 BE247575.1 EST_HUMAN 1 16840 2.82280 0.78 2.0E-35 BE323636.1 EST_HUMAN 1 16840 2.82280 0.78 2.0E-35 BE323636.1 EST_HUMAN 1 16840 2.8280 0.78 2.0E-35 BE323636.1 EST_HUMAN 1 16840 2.8280 0.65 2.0E-35 BE323636.1 EST_HUMAN 1 16840 2.8280 0.65 2.0E-35 BE323636.1 EST_HUMAN 1 16840 2.8280	Probe SEQ ID NO:			Expression Signal				Top Hit Descriptor
4 22722 36639 1.12 3.0E-35 AW003063.1 EST_HUMAN 15636 25667 1.88 2.0E-35 AW003063.1 EST_HUMAN 14943 25682 1.65 2.0E-35 AB0966.1 EST_HUMAN 14944 27683 5.73 2.0E-35 AB018413.1 NT 16066 28714 1.12 2.0E-35 AB020702.1 NT 16066 28715 1.12 2.0E-35 AB020702.1 NT 16300 28716 0.78 2.0E-35 AB020702.1 NT 16304 28280 0.78 2.0E-35 BE247575.1 EST_HUMAN 18294 31192 2.7 2.0E-35 BE32636.1 EST_HUMAN 18294 31192 2.0E-35 BE32636.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BE32636.1 EST_HUMAN 18294 31192 1.28 2.0E-35 AV722718.1 EST_HUMAN 18294 31192 1.28 2.0E-35 AV223718.1 EST_HUMAN 18294 31192 1.28 2.0E-35 AV223718.1 EST_HU	8388	1		1.42	3.0E	AF223391.1		Homo saplens calclum channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
16535 25567 1.88 2.0E-35 N89965.1 EST HUMAN 13919 25562 1.65 2.0E-35 N89965.1 EST HUMAN 14943 27683 6.73 2.0E-35 AB018413.1 NT 16066 28714 1.12 2.0E-35 AB018413.1 NT 16066 28715 1.12 2.0E-35 AB018413.1 NT 16066 28716 1.12 2.0E-35 AB020702.1 NT 16840 29279 0.78 2.0E-35 BE247575.1 EST HUMAN 17356 2.0279 0.78 2.0E-35 BE247575.1 EST HUMAN 18294 31192 2.7 2.0E-35 BE32638.1 EST HUMAN 19696 32749 0.85 2.0E-35 BE32638.1 EST HUMAN 20471 33583 0.45 2.0E-35 BE32638.1 EST HUMAN 16096 32749 0.45 2.0E-35 BE32638.1 EST HUMAN 16294 31192 1.28 2.0E-35 BE32638.1 EST HUMAN 16066 28714 1.22 2.0E-35 BE904977.1 NT <td>10074</td> <td></td> <td></td> <td>1.12</td> <td>3.0E</td> <td>AW003063.1</td> <td>EST HUMAN</td> <td>wr03a05.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2480432 3' similar to SW:POL1_HUMAN P10288 RETROVIRUS-REI ATED POI POI SPECIFIN INCOLTANDO CONTRAINS</td>	10074			1.12	3.0E	AW003063.1	EST HUMAN	wr03a05.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2480432 3' similar to SW:POL1_HUMAN P10288 RETROVIRUS-REI ATED POI POI SPECIFIN INCOLTANDO CONTRAINS
13919 26582 1.66 2.0E-35 171908.1 EST HUMAN 14943 27683 6.73 2.0E-35 AB018413.1 NT 16066 28714 1.12 2.0E-35 AB018413.1 NT 16066 28715 1.12 2.0E-35 AB020702.1 NT 16300 0.78 2.0E-35 AB247575.1 EST HUMAN 16540 29279 0.78 2.0E-35 BE247575.1 EST HUMAN 16540 29280 0.78 2.0E-35 BE247575.1 EST HUMAN 18294 31192 2.7 2.0E-35 BE32836.1 EST HUMAN 19696 32749 0.85 2.0E-35 BE832836.1 EST HUMAN 19696 32760 0.85 2.0E-35 BE832836.1 EST HUMAN 19696 32749 0.45 2.0E-35 BE832836.1 EST HUMAN 18294 31192 1.28 2.0E-35 BE832836.1 EST HUMAN 1668 2.24 2.0E-35 <t< td=""><td>106</td><td></td><td></td><td>1.88</td><td></td><td>N88965.1</td><td>EST HIMAN</td><td>K6932F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone K6932 5' similar to</td></t<>	106			1.88		N88965.1	EST HIMAN	K6932F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone K6932 5' similar to
14943 27683 5.73 2.0E-35 AB018413.1 INT 16066 28714 1.12 2.0E-35 AB020702.1 INT 16066 28714 1.12 2.0E-35 AB020702.1 INT 16640 28279 0.78 2.0E-35 BE247575.1 EST HUMAN 16640 29280 0.78 2.0E-35 BE247675.1 EST HUMAN 17356 2.7 2.0E-35 BE32477.1 EST HUMAN 19896 32749 0.85 2.0E-35 BE32836.1 EST HUMAN 19896 32749 0.85 2.0E-35 BE32836.1 EST HUMAN 20471 33583 0.45 2.0E-35 BE32836.1 EST HUMAN 20471 33584 0.45 2.0E-35 BE32836.1 EST HUMAN 16284 31192 1.28 2.0E-35 BE32836.1 EST HUMAN 16284 31120 0.45 2.0E-35 BE32836.1 EST HUMAN 16284 31120 1.72 2.0E-35 BE32417.1 INT HUMAN 16066 28714 1.72 2.0E-35 BE904978.1 EST HUMAN	1165		26582	1.65	L	T11909.1	EST HIMAN	A077F Heart Home conjune - DNA - Lange Act
16066 28714 1.12 2.0E-35 6912459 NT 16066 28715 1.12 2.0E-35 6912459 NT 16300 0.94 2.0E-35 BE24755.1 EST_HUMAN 16840 29279 0.78 2.0E-35 BE24755.1 EST_HUMAN 17356 2.57 2.0E-35 BE24755.1 EST_HUMAN 18294 31192 2.7 2.0E-35 BE33241.1 EST_HUMAN 19696 32749 0.85 2.0E-35 BE832836.1 EST_HUMAN 20471 33583 0.45 2.0E-35 BE832836.1 EST_HUMAN 20471 33584 0.45 2.0E-35 BE832836.1 EST_HUMAN 10294 31192 1.28 2.0E-35 BE832836.1 EST_HUMAN 10294 31192 1.28 2.0E-35 BE932417.1 INT 10294 31120 1.22 2.0E-35 BE904977.1 INT 10668 28714 1.72 2.0E-35 BE904977.1 INT </td <td>2215</td> <td>1</td> <td>27683</td> <td>5.73</td> <td></td> <td></td> <td>LN</td> <td>Homo sapiens mBNA for Coh?</td>	2215	1	27683	5.73			LN	Homo sapiens mBNA for Coh?
16066 28715 1.12 2.0E-35 6912459 NT 16500 0.94 2.0E-35 AB020702.1 NT 16640 29279 0.78 2.0E-35 BE247575.1 EST_HUMAN 15640 29280 0.78 2.0E-35 BE247675.1 EST_HUMAN 17356 2.57 2.0E-35 BE324767.1 EST_HUMAN 18294 31192 2.7 2.0E-35 BE32836.1 EST_HUMAN 19696 32749 0.85 2.0E-35 BE832836.1 EST_HUMAN 20471 33583 0.45 2.0E-35 BE832836.1 EST_HUMAN 20471 33584 0.45 2.0E-35 BE832836.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BE932836.1 EST_HUMAN 16284 31120 1.28 2.0E-35 BE904977.1 NT 16284 31120 1.22 2.0E-35 BE904977.1 NT 16068 28716 1.36 2.0E-35 BE904	3306		28714	1.12	2.0E-35	6912459	Į.	Homo sapiens Grb2-associated blodge 3 /VIA A A SE 22 - DAYA
16540 0.94 2.0E-35 AB020702.1 NT 16640 29279 0.78 2.0E-35 BE247575.1 EST_HUMAN 17359 2.07 2.0E-35 BE247575.1 EST_HUMAN 17359 2.0F-35 BE32636.1 EST_HUMAN 18294 31192 2.7 2.0E-35 BE32636.1 EST_HUMAN 19696 32749 0.85 2.0E-35 BE832636.1 EST_HUMAN 20471 33583 0.45 2.0E-35 BE832636.1 EST_HUMAN 20471 33584 0.45 2.0E-35 BE832636.1 EST_HUMAN 20471 33589 0.45 2.0E-35 BE832636.1 EST_HUMAN 20471 33589 0.45 2.0E-35 BE832636.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BE832636.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BE832636.1 EST_HUMAN 2388 3622 2.24 2.0E-35 BE32636.1 EST_HUMAN 24577 1.21 1.2 2.0E-35 BE904978.1 EST_HUMAN 24577 31121<	3306	- 1	28715	1.12	2.0E-35		TN	Homo saplens Grb2-associated binder 7 (KIAA6574) "PNA
16840 29279 0.78 2.0E-35 BE247575.1 EST_HUMAN 17356 29280 0.78 2.0E-35 BE247575.1 EST_HUMAN 17356 2.57 2.0E-35 BH2247575.1 EST_HUMAN 18294 31192 2.7 2.0E-35 BH32243.1 EST_HUMAN 19696 32749 0.85 2.0E-35 BE832836.1 EST_HUMAN 20471 33583 0.45 2.0E-35 BE832836.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BA722718.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BA722718.1 EST_HUMAN 20471 33594 0.45 2.0E-35 BA722718.1 EST_HUMAN 18294 31192 1.28 2.0E-35 BA722718.1 EST_HUMAN 18668 28714 1.72 2.0E-35 BE904977.1 INT 24577 31120 1.36 2.0E-35 BE904978.1 EST_HUMAN 24507 31121 1.3	3545	- 1		0.94	2.0E-35	AB020702.1	IN	Homo sapiens mRNA for KIAA0895 protein partial ofs
16840 29280 0.78 2.0E-36 BE247575.1 EST HUMAN 17356 2.57 2.0E-36 BF323477.1 EST HUMAN 18296 32749 0.95 2.0E-36 BE832636.1 EST HUMAN 18696 32750 0.95 2.0E-36 BE832636.1 EST HUMAN 20471 33563 0.45 2.0E-36 BE832636.1 EST HUMAN 20471 33564 0.45 2.0E-36 BE832636.1 EST HUMAN 20471 33569 0.45 2.0E-36 BE832636.1 EST HUMAN 16284 31182 2.24 2.0E-36 AV723718.1 EST HUMAN 16284 31182 1.28 2.0E-36 BE32636.1 EST HUMAN 16066 28714 1.72 2.0E-36 BE32477.1 NT 16066 28714 1.72 2.0E-36 BE332477.1 EST HUMAN 24577 31121 1.36 2.0E-36 BE904978.1 EST HUMAN 24577 31121 1.36 2.0E-36 BE904978.1 EST HUMAN 24500 6.98 2.0E-36 AL163210.2 NT H <td>3860</td> <td>1</td> <td>29279</td> <td>0.78</td> <td>2.0E-35</td> <td>BE247575.1</td> <td>EST_HUMAN</td> <td>TCBAP2E4328 Pediatric pre-B cell acute lymphoblastic leukemia Baylαr-HGSC project=TCBA Homo sapiens cDNA done TCBAP4328</td>	3860	1	29279	0.78	2.0E-35	BE247575.1	EST_HUMAN	TCBAP2E4328 Pediatric pre-B cell acute lymphoblastic leukemia Baylαr-HGSC project=TCBA Homo sapiens cDNA done TCBAP4328
17366 2.57 2.0E-36 H49239.1 EST HUMAN 16294 31192 2.7 2.0E-36 BF332417.1 EST HUMAN 16936 32749 0.85 2.0E-35 BE832638.1 EST HUMAN 20471 33583 0.45 2.0E-35 BE832638.1 EST HUMAN 20471 33584 0.45 2.0E-35 AV723718.1 EST HUMAN 23388 36826 2.24 2.0E-35 AV723718.1 EST HUMAN 18284 3192 1.28 2.0E-35 AV723718.1 EST HUMAN 16066 28714 1.72 2.0E-36 BF332417.1 NT 16066 28714 1.72 2.0E-36 BF332417.1 EST HUMAN 24577 31120 1.72 2.0E-36 BF30497.1 NT 24577 31120 1.36 2.0E-36 BF904978.1 EST HUMAN 16566 2.0E-36 BF904978.1 EST HUMAN EST HUMAN 16577 31121 1.36 2.0E-36 BF904978.1 EST HUMAN 12874 2.56 2.0E-36 BF904978.1 EST HUMAN	3890	16640	29280	0.78	2.0E-35		EST HUMAN	TCBAP2E4328 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo saplens cDNA ciona TCRAP4328
18294 31192 2.7 2.0E-35 BF332417.1 EST HUMAN 19696 32749 0.95 2.0E-35 BE832636.1 EST HUMAN 20471 33583 0.45 2.0E-35 BE832638.1 EST HUMAN 20471 33594 0.45 2.0E-35 AV723718.1 EST HUMAN 20471 33594 0.45 2.0E-35 AV723718.1 EST HUMAN 18294 31192 1.28 2.0E-35 K59417.1 NT 16066 28714 1.72 2.0E-35 BF32417.1 EST HUMAN 16067 28715 1.72 2.0E-36 BF32417.1 EST HUMAN 24577 31120 1.72 2.0E-36 BF304917.1 NT 24577 31120 1.72 2.0E-36 BF304978.1 EST HUMAN 24507 31121 1.36 2.0E-36 BE904978.1 EST HUMAN 1555 2.0E-36 BE904978.1 EST HUMAN 12874 2.0E-36 BE904978.1 EST HUMAN 12874 2.0E-36 AL4631949.1 EST HUMAN 12874 2.0E-36 AR4631949.1 EST HUMAN	4621			2.57	2.0E-35	Ī	FST HUMAN	VIT (901) 11 Schares fatel liner en loon ANET E. 1
19696 32749 0.95 2.0E-35 BE832836.1 EST HUMAN 20471 33583 0.45 2.0E-35 BE832836.1 EST HUMAN 20471 33594 0.45 2.0E-35 AV723718.1 EST HUMAN 20471 33594 0.45 2.0E-35 AV723718.1 EST HUMAN 18294 31192 1.28 2.0E-35 K59417.1 NT 18096 28714 1.72 2.0E-35 BF32417.1 EST HUMAN 16066 28714 1.72 2.0E-35 BF32417.1 EST HUMAN 24577 31120 1.72 2.0E-35 BF30497.1 EST HUMAN 24577 31120 1.36 2.0E-35 BE904978.1 EST HUMAN 24500 5.98 2.0E-35 BE904978.1 EST HUMAN 12874 25456 1.56 2.0E-35 BE904978.1 EST HUMAN 12874 25496 6.81 1.0E-35 AR31949.1 EST HUMAN 12874 25497 6.81 1.0E-35 AR31949.1 EST HUMAN 13609 25667 1.0E-35 AR31949.1 EST HUMAN	5495		31192	2.7	2.0E-35		EST HUMAN	OVO-BT0701-210400-100 PA BT0201 Dome 11 PT CO TOUR SECTION CODE IMAGE:274079 6
19696 32760 0.85 2.0E-35 BE832838.1 EST_HUMAN 20471 33583 0.45 2.0E-35 AV723718.1 EST_HUMAN 20471 33594 0.45 2.0E-35 AV723718.1 EST_HUMAN 18294 31192 1.28 2.0E-35 K59417.1 NT 16068 28714 1.72 2.0E-35 BF322417.1 EST_HUMAN 16069 28714 1.72 2.0E-35 BF322417.1 EST_HUMAN 24577 31120 1.72 2.0E-35 BF32417.1 EST_HUMAN 24577 31120 1.36 2.0E-35 BE904978.1 EST_HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST_HUMAN 24500 5.98 2.0E-35 BE904978.1 EST_HUMAN 12874 25456 6.81 1.0E-35 AL18310.2 NT 12874 25496 6.81 1.0E-35 AR31949.1 EST_HUMAN 13609 26165 1.0E-35 AR31949.1 EST_HUMAN 13609 26165 1.0E-35 AR31949.1 EST_HUMAN 13609	7004		32749	0.95	2.0E-35		EST HUMAN	CM2-MT0125-280700-397-Gn2-MT0125-Discontinuing September CDNA
20471 33563 0.45 2.0E-35 AVT23718.1 EST_HUMAN 20471 33594 0.45 2.0E-35 AVT23718.1 EST_HUMAN 12388 36626 2.24 2.0E-35 X59417.1 NT 18294 31192 1.28 2.0E-35 BF322417.1 EST_HUMAN 16068 28714 1.72 2.0E-35 BF322417.1 EST_HUMAN 16069 28714 1.72 2.0E-35 BF32417.1 EST_HUMAN 24577 31120 1.36 2.0E-35 BF904878.1 EST_HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST_HUMAN 24500 5.98 2.0E-35 BE904978.1 EST_HUMAN 12874 25667 1.56 2.0E-35 BE904978.1 EST_HUMAN 12874 25496 6.81 1.0E-35 A4831949.1 EST_HUMAN 13609 26165 1.0E-35 A4831949.1 EST_HUMAN 13609 26165 1.0E-35 A48319473.1 EST_HUMAN	7007	- 1	32750	0.95	2.0E-35		EST HUMAN	CM2-MT0125-280700-297-(902 MT0125 Home september 2014A
20471 33594 0.45 2.0E-35 AV723718.1 EST HUMAN 23388 36626 2.24 2.0E-35 K59417.1 NT 18294 31192 1.28 2.0E-35 BF322417.1 EST HUMAN 16068 28714 1.72 2.0E-35 BF322417.1 EST HUMAN 24577 31120 1.72 2.0E-35 BF904978.1 EST HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 24507 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 16535 25567 1.56 2.0E-35 AL163210.2 NT 12874 25496 6.81 1.0E-35 AR631949.1 EST HUMAN 12874 25497 6.81 1.0E-35 AR631949.1 EST HUMAN 13609 26165 1.0E-35 AR939473.1 EST HUMAN	7775	- 1	33583	0.45	2.0E-35,		EST HUMAN	AV723718 HTB Home sapiens chive Libra Libra Libra Walls and
23388 36626 2.24 2.05.36 K59417.1 NT 18294 31192 1.28 2.05.35 BF332417.1 EST_HUMAN 16068 28714 1.72 2.0E.35 BF332417.1 EST_HUMAN 24577 31120 1.72 2.0E.35 BF90459 NT EST_HUMAN 24577 31120 1.36 2.0E.35 BE904978.1 EST_HUMAN 24677 31121 1.36 2.0E.35 BE904978.1 EST_HUMAN 24600 5.99 2.0E.35 AL163210.2 NT 12874 25497 6.81 1.0E.35 AA631949.1 EST_HUMAN 12874 25497 6.81 1.0E.35 AA631949.1 EST_HUMAN 13609 26165 1.0E.35 AA631949.1 EST_HUMAN 13609 26165 1.0E.35 AA631949.1 EST_HUMAN	7776	- 1	33594	0.45	2.0E-35,		EST HUMAN	AV723718 HTB Homo seriens cDNA clone HTBAVA10 &
18284 31192 1.28 2.0E-35 BF322417.1 EST HUMAN 16068 28714 1.72 2.0E-35 BF322417.1 EST HUMAN 24577 31120 1.72 2.0E-35 BE904978.1 EST HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 24507 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 1550 5.99 2.0E-35 AL163210.2 NT 1551 1.56 2.0E-35 AL163210.2 NT 12874 25495 6.81 1.0E-35 AR31949.1 EST HUMAN 12874 25497 6.81 1.0E-35 AR31949.1 EST HUMAN 13509 26165 1.0E-35 AR339473.1 EST HUMAN	10697	23388	36626	2.24	2.0E-35),		L L	H.sapiens PROS-27 mRNA
16066 28714 1.72 2.0E-35 6912459 INT 16066 28716 1.72 2.0E-35 6912459 INT 24577 31120 1.36 2.0E-35 BE904978.1 EST HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 24500 5.98 2.0E-35 AL163210.2 NT 15256 1.56 2.0E-35 AR631949.1 EST HUMAN 12874 25496 6.81 1.0E-35 AR631949.1 EST HUMAN 13609 28165 1.0E-35 AR631949.1 EST HUMAN 13609 28165 1.0E-35 AR9389473.1 EST HUMAN	11817	18294	31192	1.28	2.0E-35		EST_HUMAN	QV0-BT0701-210400-199-b04 BT0701 Homo septems cDNA
16066 28716 1.72 2.0E-35 6912459 INT 24577 31120 1.36 2.0E-35 BE904978.1 EST_HUMAN 24577 31121 1.36 2.0E-35 BE904978.1 EST_HUMAN 24500 5.98 2.0E-35 AL163210.2 NT 15535 2.5567 1.56 2.0E-35 AR31949.1 EST_HUMAN 12874 25495 6.81 1.0E-35 AA631949.1 EST_HUMAN 13609 28165 1.66.31 AA631949.1 EST_HUMAN 13609 28165 1.0E-35 AA831949.1 EST_HUMAN	11889	16068	28714	1.72	2.0E-35	6912459	4	Homo saplens Grb2-associated binder 2 /KIAA0574 mBAIA
24577 31120 1.36 2.0E-36 BE904978.1 EST_HUMAN 24577 31121 1.36 2.0E-36 BE904978.1 EST_HUMAN 24900 5.98 2.0E-36 AL163210.2 NT 15536 2.5667 1.56 2.0E-36 N89965.1 EST_HUMAN 12874 25496 6.81 1.0E-36 AA631949.1 EST_HUMAN 13609 28165 1.66.31 AA631949.1 EST_HUMAN	1889	16066	28715	1.72	2.0E-35	6912459	5	Homo saplens Grb2-associated hinder 2 (KLAAG74)
24577 31121 1.36 2.0E-35 BE904978.1 EST HUMAN 24900 6.98 2.0E-35 AL163210.2 NT 15535 2.5667 1.56 2.0E-35 N88966.1 EST HUMAN 12874 25496 6.81 1.0E-35 AA631949.1 EST HUMAN 12874 25497 6.81 1.0E-36 AA631949.1 EST HUMAN 13609 28165 1.96 1.0E-35 AAV389473.1 EST HUMAN	12082	24577	31120	1.36	2.0E-35		T HUMAN	601498774F1 NIH MGC 70 Home sepiens e 2000 i i iliniya
24900 6.98 2.0E-35 AL163210.2 NT 15535 25667 1.56 2.0E-35 N89965.1 EST_HUMAN 12874 25496 6.81 1.0E-35 AA631949.1 EST_HUMAN 12874 25497 6.81 1.0E-36 AA631949.1 EST_HUMAN 13609 28165 1.96 1.0E-35 AW389473.1 EST_HUMAN	12062	24577	31121	1.36			Г	601496774F1 NIH MGC 70 Homo saplens cDNA clone IVA CE: 3006000 51
15636 25667 1.58 2.0E-35 N89965.1 EST_HUMAN 12874 25496 6.81 1.0E-35 AA631949.1 EST_HUMAN 12874 25497 6.81 1.0E-36 AA631949.1 EST_HUMAN 13609 28165 1.9.5 1.0E-35 AW389473.1 EST_HUMAN	12572	24900		6.98				Homo saplens chromosome 21 segment HS21C010
12874 25496 6.81 1.0E-35 AA631949.1 EST HUMAN 12874 25497 6.81 1.0E-35 AA631949.1 EST HUMAN 13609 28165 1.9E-3 AW389473.1 EST HUMAN	12689	15535	25567	1.58	2.0E-35 N			K6832F Human fetal heart, Lambda ZAP Express Homo saplens cDNA clone K6932 6' similar to REPETITIVE FI EMENT
12874 25497 6.81 1.0E-35 AA631949-1 EST HUMAN 13509 28165 19.5 1.0E-35 AW389473-1 EST HUMAN	45	12874	25496	6.81	1.0E-35 A		T	finitional genomic DNA specific CDNA library
13509 28165 19.5 1.0E-35 AW389473.1 EST_HUMAN	45	12874	25497	. 6.81	1.0E-35		T	finfort Beginnel cenomic DNA specific cDNA library nome sapiens cDNA clone CR12-1
	736	13509	28165	19.5	1.0E-35		Т	112-ST0162-13109-00-412-ST0162-Home carlana china china

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	П					Т	П		Mus musculus activin receptor interacting protein 1 (Arip1-pending), mRNA	Mus musculus activin receptor interacting protein 1 (Arip1-pending), mRNA	Homo sapiens chromatin assembly factor 1, subunit B (p60) (CHAF1B), mKNA	Homo sapiens mRNA for KIAA1279 protein, partial cds			П		Г	┑	Homo sapiens mRNA for KIAA1057 protein, pardai cas	П	ヿ	Homo sapiens fibulin 1 (FBLN1), mKNA	П	IMAN 601584833F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3438983 5		ヿ	Homo sapiens C-terminal binding protein 2 (CLBP2) mKNA	Human carcinoembryonic antigen gene family member 12 (Cow12) gate, explising the Color (Cow12)	Human carcinoembryonic antigen gene family member 12 (CGM12) gene, excits Land DIV
Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST HUMAN	Ι	EST_HUMAN	EST_HUMAN	NT	LΝ	LΝ	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST HUMAN	Ę	N	EST_HUMAN	뉟	ΙΝ	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN.	뉟	뉟
Top Hit Acession No.	1.0E-35 AW389473.1	35 T87947.1	7705994 NT	35 BE350127.1	35 RE350127 1	B006030 NT	.35 AV650422.1		7656905 NT	7656905 NT	11526236 NT	1.0E-35 AB033105.1	11418002 NT	-35 AU158595.1	-35 AU158595.1	35 BF589594.1		-35 BF589594.1	-35 AB028980.1	-35 AB028980.1	-35 AI525119.1		11418110 NT	-35 BE792832.1	8.0E-36 AA348480.1	AW8575	4557498 NT	7.0E-36 U06672.1	7.0E-36 U06672.1
Most Similar (Top) Hit BLAST E Value	1.0E-35	1.0E-35	1.0E-35	1.0E-35	4.0F_35	105.35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35		1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	8.0E-36	7.0E-36	7.0E-36	7.0E-36	7.05-36
Expression Signal	19.5	1.3	1.68	8.	2	2.5	3.3	3.3	3.83	3.93	1.41	0.86	1.18	2.16	2.16	0.7		0.7	1.46	1.46	1.91	1.37	1.63	2.13	0.56	1.1	3.84	5.92	
ORF SEQ ID NO:	26166		27996	28217	9535	285/0	28570	28571	29756	29757	30934	33141	33311	35297	35298	36352		36353	37680	37681					34983	28344		33327	33328
Exan SEQ ID NO:	13509	13658	15258	15475	76778	2652	15924	15924	17125	17125	18222	20063	20211	25125	25125	23123		23123	24349	24349		25313	24613	24837	21817	1	15881		Ш
Probe SEQ ID NO:	735	88	2544	2770		2/72	3161	3161	4388	4388	5423	7383	7541	9442	B442	10477		10477	11758	11758	11768	11917	12121	12471	9129	2331	3116	7554	7664

Page 270 of 536 Table 4 Single Exon Pmhae Exonation 1

Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sepiens niniurin 2 (NINJ2), mRNA	Hamo saolens TCI 6 gene exon 12	UI-H-BW1-anv-c-12-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA done IMAGF:3083542 31	193506.x1 Soares NSF F8 9W OT PA P S1 Homo sapiens cDNA clone INAGE:2126195 3' cimilar to gb:M11949 PANCREATIC SECRETORY TRYPSIN INHIBITOR PRECLIBSOR (HINAN)	ho06h02.xt NCI_CGAP_Co14 Homo saplens cDNA clone IMAGE:3036627 3' similar to SW:IMA2_HUMAN P62292 IMPORTIN ALPHA-2 SUBUNIT	Homo saplens syncytin precursor, mRNA, complete cde	C16927 Clontech human aorta bolyA+ mRNA (#8572) Home sanians cDNA clone CEN E3EC14 E	#9509.x1 NC_CGAP_CL1 Homo sapiens cDNA clone IMAGE:2107024 3' similar to contains MER9.b2	Homo sablans Xd bseudosumal region: segment 1/2	601285667F1 NIH MGC 44 Homo sepiens cDNA clone (MAGE:3807289 5'	Homo sapiens chromosome 21 segment HS21C009	Homo saplens APIG-like 1 (APISL1), mRNA	Homo sapiens API6-like 1 (API5L1), mRNA	Homo sapiens N-ethylmaleimide-sensitive factor (NSF), mRNA	Homo sapiens Xq pseudoautosomal region: segment 1/2	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	PM3-BN0176-100400-001-g04 BN0176 Homo sapiens cDNA	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE;	601298674F1 NIH MGC 19 Homo septemb cDNA clone IMAGE 3638388 F1	2820020.5prime NIH MGC 7 Homo seplens cDNA clone (MAGE-2820020 F)	601282266F1 NIH MGC 44 Homo sepiens cDNA clone IMAGE:36041AR F	601282286F1 NIH MGC 44 Homo sapiens cDNA clone IMAGE:3604188 F	W19f05.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE-130713 51	Homo saplens a disintegrin and metalloprotelnase domain 22 (ADAM22), transport variant 3 mRNA	Human platelet Glycoprotein (Ib (GPIIb) gene, exons 2-29	Homo saplens DNA for amyloid precursor protein, complete cds	Homo sapiens DNA for amyloid precursor protein, complete cds	zu89c10.r1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:743250 67	Homo sapiens nuclear factor of activated T-cells, cytoplasmic 2 (NFATC2), mRNA	AV763929 TP Homo saplens cDNA clone TPGABH01 6'
gle Exon Prob	Top Hit Database Source			EST_HUMAN	EST_HUMAN		Т	T HUMAN		Т	T HUMAN	Г				Į.		EST_HUMAN F	A TORGASIWS	Г	Г	Г	П	Г				H L	T_HUMAN		EST_HUMAN A
Sing	Top Hit Acession No.	7706622 NT	6.0E-36 AB035346.1	6.0E-36 BF515101.1	6.0E-36 A1435169.1	6.0E-36 AW780143.1		Π	6.0E-36 Al380499.1	Ī	5.0E-38 BE388436.1	Γ	5729729 NT	5729729 NT	11079227 NT	6.0E-36 AJ271735.1	11417862 NT	-36 BE010038.1		4.0E-36 BE382574.1		-36 BE389299.1 E	-36 BE389299.1	-36 R64023.1	11497041 NT				4.0E-36 AA400370.1	516	4.0E-36 AV753829.1 E
	Most Similar (Top) Hit BLAST E Value	6.0E-36	6.0E-36	6.0E-36	6.0E-36	6.0E-36	6.0E-36	6.0E-36	6.0E-36	5.0E-38/	6.0E-38	5.0E-36	5.0E-36	5.0E-38	5.0E-36	6.0E-36	5.0E-36	4.0E-36	4.0E-36 F	4.0E-38 E	4.0E-38	4.0E-36	4.0E-36 B	4.0E-36 R	4.0E-36	4.0E-36 M33320.1	4.0E-36 D87675.1	4.0E-36 D87675.1	4.0E-36	4.0E-38	4.0E-36 A
	Expression Signal	2	5.58	0.71	3.54	3.57	2.33	0.51	3.11	10.74	5.75	1.45	2.15	2.15	0.61	3.53	3.45	1.69	1.03	1.61	2.13	0.82	0.82	0.84	2.33	1.63	1.62	1.62	2.84	2.09	7.3
	ORF SEQ ID NO:	27456		23023	30682	32766	34385		37443	25592	28202	28991	30104	30105	33464	26592	31103	26619	26856	27074		28763	28764		31707	33326	34285	34286	36835	1	
	Exan SEQ ID NO:			16383	18054	19701	ı	22773	24136	12949	15460	16352	17488	17468	20350	12949	24650	13965	14170	14388	14947	16108	16108	18425	18746	20223	21145	21145	23589	24655	25199
	Probe SEQ ID NO:	1998	2418	3830	6248	7009	8550	10126	11536	134	2756	3599	4736	4736	7886	11887	12168	1203	1423	1840	2219	3349	3349	9829	5964	7683	8453	8453	10909	12183	12227

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						שם ביצימו בומר	Single Exoli Flores Expressed in plant
Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
12798	25047		1.44	4.0E-36	36 D25217.2	NT	Homo saplens mRNA for KIAA0027 protein, partial cds
88		26100		3.0E-36	36 AF099810.1	LN	Homo sapiens neurexin III-alpha gene, partial cds
1484	14231	26917	1.32	3.0E-36	36 AF110239.1	NT	Homo sapiens calcium/calmodulin-stimulated cyclic nucleotide phosphodiesterase (PDE1A) gene, partial ods
484	14231	26918	1.32	3.0E-36	36 AF110239.1	FN	Homo sapiens calcium/calmodulin-stimulated cyclic nucleotide phosphodiesterase (PDE1A) gene, parttal cds
2297				3.0E-36		NT	Homo sapiens KIAA0952 protein (KIAA0952), mRNA
4467	17203	<u> </u>	98.9	3.0E-36	10181139 NT	TN	Mus musculus junctophilin 1 (Jp1-pending), mRNA
11050	23720	36991	1.59	3.0E-36	36 BF035327.1	EST HUMAN	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5
3167	15930			2.0E-36	36 BE259267.1	EST HUMAN	601106343F1 NIH_MGC_16 Homo sapiens cLINA cione IMAGE:3342/00 3
4904	17831	30246	5.45	2.0E-36	36 AW880376.1	EST HUMAN	QV0-OT0030-240300-174-h04 OT0030 Hamo sapiens CDNA
5398	18198	30892		2.0E-	36 AF267747.1	NT	Mus musculus p47-phax gene, complete cds
222	18550	31471	3.95	2.0E-	36 T08756.1	EST_HUMAN	EST06648 Infant Brain, Bento Soares Homo sapiens cUNA clone HIBBU28 3 end
6481	!	32248	1	2.0E-36	36 T69629.1	EST_HUMAN	yc44a07.r1 Stratagene Iver (#837224) Homo saplens cDINA cione IMAGE:83508 5
9288	21855	35126		2.0E-	36 BF512794.1	EST_HUMAN	UI-H-BW1-amu-a-11-0-UI.s1 NO CGAP Sub7 Homo sepiens cDNA cione IMAGE: 30/1132.3
9449	21999	35172	0.79	2.0E-		Ä	Homo sapiens ubiquitin specific protease 13 (isopeptidase 1-3) (USP13) mRNA
9448	21999	35173	62'0	2.0E-	4507848 NT	L	Homo sapiens ubiquitin specific protease 13 (isopeptidase 1-3) (USP13) mRNA
865	13634	26304	1.81	1.0E-36	36 BE409310.1	EST_HUMAN	601300938F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3635480 5
2141	14871	27603	1	1.0E-36	36 BE146523.1	EST HUMAN	RC1-HT0217-131199-021-h07 HT0217 Homo sapiens CDNA
2141	14871	27604	-	1.0E-36	36 BE146523.1	EST_HUMAN	RC1-HT0217-131199-021-h07 HT0217 Homo saplens cDNA
2199	ı	27664	1.36	1.0E-	36 BF673761.1	EST_HUMAN	602136493F1 NIH_MGC_83 Homo sapiens cDNA cione IMAGE: 427/2886 5
3339	16099		1.16	1.0E	36 AF156962.1	Z	Homo sapiens human endogenous retrovirus W proC6-19 protease (pro) gene, paraal cos
5810	18599	31527	1.29	1.0E-36	4827064 NT	۲	Homo saplens zinc finger protein 147 (estrogen-responsive linger protein) (ZNr147) mKNA
			,	7	20 A 10 6 774 4 4	H TAN	wb37c12x1 NCI_CGAP_GC6 Homo sapiens cDNA cione IMAGE:x307662 5 similar to contatus Aurireneefitive element:
OROG	10000		ř	2	1.1.1		yg36g10.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:34529 6' similar to
6296	19069	32052	12	1.0E-38	36 R25012.1	EST_HUMAN	SP:CAHP_HUMAN P35219 CARBONIC ANHYDRASE-RELATED PROTEIN;
900	1	_	:	10.	36 R25012.1	EST HUMAN	yg36g10.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:34529 6' similar to SP:CAHP_HUMAN P35219 CARBONIC ANHYDRASE-RELATED PROTEIN;
AED 2	ı			199	36 AL 120542.1	EST HUMAN	DKFZp781A229_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761A229 5'
72.06	L			19	11426108 NT	Į.	Homo saplens a disintegrin and metalloproteinase domain 11 (ADAM11), mRNA
220	1			io t		Ę	Homo sapiens a disintegrin and metalloproteinase domain 11 (ADAM11), mRNA
250	20002			2 0	AA1480	EST HUMAN	2051a12.r1 Stratagene endothelial cell 937223 Homo sapiens cDNA clone IMAGE: 590398 61
3		١		3 5	A A 4 4 9 0 2 4 4	EST LIMAN	7051a12 rt Strategene endothelial cell 937223 Homo sapiens cDNA clone IMAGE:590398 5
7860	20555	33680	5.13	1.0E	36 AA148U34.1	ביילוויוסון וכם	

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	nc60e08.r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:745670	nc60e08.r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:745670	AU141688 THYRO1 Homo saplens cDNA clone THYRO1001033 5/	AU141688 THYRO1 Homo sapiens cDNA clone THYRO1001033 5'	xe82b07.x1 NCI_CGAP_Brn35 Homo saplens cDNA clone IMAGE:2814387 3'	QV3-NN1023-010600-199-h01 NN1023 Homo saplens cDNA	RC3-CT0279-040500-017-a10 GT0279 Homo saplens cDNA	RC3-CT0279-040500-017-a10 CT0279 Homo saplens cDNA	CM3-NN0061-140400-147-h12 NN0081 Homo sapiens cDNA	UI-HF-BNO-ale-c-03-0-UI.r1 NIH_MGC 50 Homo sablens cDNA clone IMAGE:3079277 6	RC-BT091-210199-110 BT091 Homo saplens cDNA	RC-BT091-210199-110 BT091 Homo sapiens cDNA	Homo saplens Ran GTPase activating protein 1 (RANGAP1), mRNA	Homo sapiens chromosome 21 segment HS21C013	Homo sapiens Sad1 unc-84 domain protein 2 (SUN2) mRNA, partial cds	ws80b07.x1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:25042453	ws80b07x1 NCI_CGAP_Co3 Homo saplens cDNA clone IMAGE:26042453'	73D4 Human retina cDNA Tsp509I-cleaved sublibrary Homo sapiens cDNA not directional	Homo sapiens chimerin (chimaerin) 2 (CHN2) mRNA	CM0-UT0003-050800-503-d09 UT0003 Homo saplens cDNA	ht08g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146266 3' similar to contains MER29.b3	https://www.comming.com/sepiens.com/sepien	MER29 repetitive element;	RC1-CN0008-210100-012-e09_1 CN0008 Homo sapiens cDNA	H.sapiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RINGB, 9, 13 and 14 genes	DKFZp434E0422_r1 434 (synonym: htes3) Hamo sapiens cDNA clone DKFZp434E0422 6'	Homo sapiens jun dimertzation protein gene, partial cds: cfos gene, completo cds: end unknown nene	Homo saplens tun dimerization protein gene partial cyts often gene powentate cyts and untowns ages	wk26b11.x1 NCI_CGAP_Bm25 Homo saplens cDNA clone IMAGE:2413341 3' similar to contains PTR5.12 PTR5.12
gle Exon Probe	Top Hit Database Source	EST_HUMAN	EST_HUMAN r	Γ		EST_HUMAN	EST_HUMAN (EST_HUMAN F	EST HUMAN	г	EST_HUMAN		Т		LN LN	LN	EST HUMAN V	EST_HUMAN v	П		T_HUMAN	T NAME IN	Т	EST_HUMAN N	EST_HUMAN R	TN P	L HUMAN	TN	I	T_HUMAN
Sin	Top Hit Acession No.	-36 AA420487.1	1.0E-36 AA420467.1	1.0E-36 AU141688.1	1.0E-36 AU141688.1	1.0E-36 AW103658.1	1.0E-36 BF364169.1	1.0E-36 AW855868,1	1.0E-36 AW855868.1	-36 AW897636.1	-36 AW504143.1		-36 Al905536.1	11418177 NT			Γ		-37 W22618.1	4757979 NT	37 BE698077.1	37 BE380427 4	T		37 AW840840.1	37 X87344.1	-	37 AF111167.2	37 AF111167.2	
	Most Similar (Top) Hit BLAST E Value	1.0E-38	1.0E-36	1.0E-36	1.0E-38	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-36	1.0E-38	9.0E-37	9.0E-37	9.0E-37	8.0E-37	8.0E-37	8 0E.37		8.0E-37	8.0E-37	8.0E-37	7.0E-37	7.0E-37	7.0E-37	
	Expression Signal	1.34	1.34	0.61	0.61	2.71	3.88	0.56	0.56	3.3	4.17	1.45	1.45	3.81	3.03	3.23	2,12	2.12	1.35	66.0	1.58	3 75		3.75	8.24	822	3.03	0.97	0.97	8.69
	ORF SEQ ID NO:	33770	33771	33902	33903	34762	35878	36086	36087	36795	37347	37302	37303				33042	33043		28765		31451		31452	31507	33604		27179	27180	36585
	SEQ ID. NO:	20647				21618	22682	22874	22874	23547	24044	23999	23999	24575	24855	25011	18965	19965	24733	16109	17977	18530		18530	18578	20479	14011	14480	14480	23348
	Probe SEQ ID NO:	7952	7952	8079	8079	8927	10014	10226	10228	10867	11354	11393	11393	12060	12501	12747	7281	7281	12309	3350	5168	5738		5738	5787	7184	1282	1738	1738	10657

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	· · · ·	Homo sapiens protocadherin alpha 10 alternate isoform (PCDH-alpha10) mRNA, complete cds			П		Homo saplens glycine C-acetyftransferase (2-amino-3-ketobutyrate-CoA ligase) (GCAT), mRNA						IAN DKFZp434L2418_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434L2418		IAN DKFZp547G067_1 547 (synonym: hfbr1) Homo sepiens cDNA done DKFZp547G067 5	at34c05.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:2373896 3' similar to TR:Q13537 AN Q13537 SIMILAR TO POGO ELEMENT.;	Homo sepiens mRNA for AML1, complete cds	Homo sapiens mRNA for AML1, complete cds	Г		Homo sapiens chromosome 21 segment HS21C047	Homo sapiens cytochrome P450, subfamily XXVIIA (sterold 27-hydroxylase, carebrotendinous	xanthomatosis), polypeptide 1 (CYPZ/A1b) mRNA							Homo saplens J domain containing protein 1 isoform b (JDP1) mRNA, complete cds	Homo saplens pescadillo (zebrafish) homdog 1, containing BRCT domain (PES1), mRNA	Homo sapiens chromosome 21 segment HS21C081
Top Hit Database Source	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	NT		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	N.	Į.	EST_HUMAN	EST_HUMAN	TN		ž	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΙN	M	님
Top Hit Acesslon No.	7.0E-37 AI536702.1	6.0E-37 AF169689.1	6.0E-37 AF202723.1	5.0E-37 AA307123.1	5.0E-37 AA307123.1	5.0E-37 AV750211.1	7857117 NT	5.0E-37 AF149773.1	4.0E-37 AA702794.1	4.0E-37 AW794502.1	AA843806.1	3.0E-37 AL048956.1	3.0E-37 AL048956.1	3.0E-37 AW961150.1	3.0E-37 AL138274.1	3.0E-37 AI749952.1	D89790.1	D89790.1	AU131202.1	2.0E-37 AU131202.1	2.0E-37 AL163247.2		4503210 NT	2.0E-37 AL163284.2	BF035327.1	2.0E-37 AA346720.1	2.0E-37 BE537764.1	2.0E-37 BE537764.1	BF204032.1	2.0E-37 AF176013.1	2.0E-37 11417972 NT	AL163281.2
Most Similar (Top) Hit BLAST E Value	7.0E-37	6.0E-37	6.0E-37	5.0E-37	5.0E-37	5.0E-37	5.0E-37	5.0E-37	4.0E-37	4.0E-37	4.0E-37	3.0E-37	3.0E-37	3.0E-37	3.0E-37	3.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37		2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	2.0E-37	1.0E-37
Expression Signal	2.25	1.34	2.94	3.9	3.9	6.0	4	6.86	2.12	0.61	0.74	3.2	3.2	3.15	0.92	0.71	0.68	0.68	2.64	2.64	1.67		4.78	0.93	99.0	3.46	0.48	0.48	2.88	11.22	3.54	4.93
ORF SEQ ID NO:	36729	34168		31744	31745	34490			27877	31945		27472			31494	33221		25843					29261			32333	33720			37459		27546
Exan SEQ ID NO:	23494	21031	24929	l	18783	l.		24572	15144	1		14745		15731	18565		L						16623	17693	l		ı	1				14813
Probe SEQ ID NO:	10811	8338	12624	6002	6002	8654	10837	12055	2423	6194	9226	2010	2010	2965	5774	7455	372	372	1058	1058	1956		3873	4968	5304	6561	7895	7895	7937	11549	12784	2081

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Single Exon Propes Expressed in Diam	Top Hit Descriptar	SSU72 PROTEIN	SSU72 PROTEIN	601157633F1 NIH_MGC_21 Homo septens cDNA clone IMAGE:3504272 5'	Homo saplens chromosome 21 segment HS21C100	xw04d01x1 NCI_CGAP_Bm53 Hamo sapiens aDNA clane IMAGE:2827009 3'	CM3-FT0181-140700-241-f07 FT0181 Homo saplens cDNA	yv88b04,r1 Scares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:249775 5'	yv88b04.r1 Scares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:249775 5'	Homo saplens chromosome 21 segment HS21C048	Homo sapiens chromosome 12 open reading frame 3 (C12ORF3), mRNA	Homo sapiens chromosome 21 segment HS21C048	Homo sapiens SMT3 (suppressor of mif two 3, yeast) homolog 2 (SMT3H2), mRNA	zw30d01.r1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:770785 5' similar to SW:MA12_RABIT P45701 MANNOSYL-OLIGOSACCHARIDE ALPHA-1,2-MANNOSIDASE;	ZW30d01.rl Sogres ovary fumor NbHOT Homo sapiens cDNA clone IMAGE:770785 5' similar to SW4.844.9 DADIT DAS704 MANNOSYI JOLIGOSACCHARIDE ALPHA-1.2-MANNOSIDASE	Addang of Spares fetal heart NhHH19W Home spiens CDNA clone IMAGE:345684 5		Homo sapiens michal for ankym b (440 kDa)	Homo sapiens mRNA for ankyrin B (440 kDa)	AV721103 HTB Homo sepiens cDNA clone HTBARH11 5'	MR3-HT0487-150200-113-g01 HT0487 Home sapiens cDNA	HSC18F031 normalized infant brain cDNA Homo saplens cDNA clone c-18f03	Homo sapiens orphan G protein-coupled receptor HG20 (HG20) mRNA, complete cds	hu09g02.x1 NCI_CGAP_Lu24 Homo sapiens cDNA done IMAGE:3166130 3' similar to TR:002710 002710 GAG POLYPROTEIN ;	Homo sapiens mRNA for KIAA0145 protein, partial cds	no34g03.s1 NCI_CGAP_Pr23 Homo sapiens cDNA done INAGE:1102812 3' similar to TR:E212316 E212316 NADP DEPENDENT LEUKOTREINE B4 12-HYDROXYDEHYDROGENASE ;	no34g03.s1 NCI_CGAP_Pr23 Homo sepiens cDNA clone IMAGE:1102812 3' similar to TR:E212316 E212316 NADP DEPENDENT LEUKOTREINE B4 12-HYDROXYDEHYDROGENASE ;	QV2-HT0698-080800-293-e05 HT0698 Homo saplens cDNA	Homo sapiens leucine-rich repeat-containing G protein-coupled receptor 6 (LGR6) mRNA, partial cds
le Exon Propes	Top Hit Database Source	SWISSPROT SS	SWISSPROT SS	EST HUMAN 60	NT Ho	П			T_HUMAN	NT		NT Ho		EST_HUMAN SV	WS NAME TO	Т	NAMOL		П		EST_HUMAN MF	T_HUMAN		EST HUMAN GA	Г	EST HUMAN E2		EST_HUMAN Q	¥ E
Buis	Top Hit Acession No.			8 BE279301.1 E	8 AL 163300.2	8 AW302461.1 E	8 BF373664.1 E			3.0E-38 AL163248.2	11435947 NT	8 AL163248.2	TN 202069	8 AA437353.1											2.0E-38 D63479.2				2.0E-38 AF190501.1
	Most Similar (Top) Hit BLAST E Value	3.0E-38 P53538	3.0E-38	3.0E-38	3.0E-38	3.0E-38	3.0E-38 €	3.0E-38 H85494.1	3.0E-38 H85494.1	3.0E-38	3.0E-38	2.0E-38/	2.0E-38	2.0E-38	L											ļ	2.0E-38,	2.0E-38	
	Expression Signal	1.76	1.76	1.47	8.11	0.56	8.26	2.1	2.1	2.24	1.65	4.1	2.99	2.21		17.7	1.40	0.69	69.0	1.46	4.38	0.56	2.04	1.06	1.71	1.37	1.37	5.79	3.52
	ORF SEQ ID NO:	29224	29226		32430	32800	33252	l			30488		26781	<u>L</u>				31339	31340			34631	L		36212			l	37557
	Exan SEQ ID NO:	16587	l.	l_	Ľ	L	L	<u></u>	21240	L.	17896	12878				- 1	ı		18427	l	L	21485	L	l	22983	.l	1.	1	24235
	Probe SEQ ID NO:	3836	3836	4574	6655	7144	7488	8548	8548	9872	12630	49	1358	1641		1641	2408	5632	5632	7619	8382	8783	8864	9121	10346	11200	11200	11472	11638

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			T	Τ	Τ	Т	Τ		Т	T	T	Τ	Γ	Τ	Τ	Γ	Ť	Τ	[8	Τ	Γ	Т	Т	Γ	8	Т	F	T -	Т
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens leucine-rich repeat-containing G protein-coupled receptor 8 (LGR8) mRNA, partial cds	AV726988 HTC Homo sapiens cDNA clone HTCAXH07 5	Hamo sapiens gene for kinesin-like protein, complete cds	CHR220580 Chromosome 22 exon Homo sapiens cDNA clone C22_788 6'	E1 beta=pyruvate dehydrogenase beta (promoter) [human, placenta, Genomic, 1280 nt]	Homo sapiens sulfotransferase-related protein (SULTX3), mRNA	zu62b02.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:742639 6' similar to contains element MER19 repetitive element;	Homo sapiens guanine nucleotide binding protein-like 1 (GNL1), mRNA	Homo saplens KIAA0173 gene product (KIAA0173), mRNA	Homo sapiens cyclin K (CCNK) gene, exon 7	Homo sapiens chromosome 21 segment HS21C003	Homo sapiens chromosome 21 segment HS21C003	Homo sapiens hypothetical protein FLJ10600 (FLJ10600), mRNA	Mus musculus otogelin (Otog), mRNA	Mus musculus atagelin (Otag), mRNA	Homo sapiens mRNA for KIAA0612 protein, partial cds	Homo sapiens hypothetical protein FLJ10600 (FLJ10600), mRNA	ht09g01.x1 NCI_CGAP_Kld13 Homo saplens cDNA clone IMAGE:3146256 3' similer to contains MER29.b3 MER29 repetitive element:	J/99b08.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:30486 5'	Homo sapiens KIAA0426 gene product (KIAA0426), mRNA	Homo saplens chromosome 21 segment HS21C084	Homo saplens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 16kD (ATP6C) mRNA	Homo saplens estrogen receptor-binding fragment-associated gene 9 (EBAG9) mRNA	wh83f10.x1 NCI_CGAP_Kid11 Homo sapiens oDNA clone IMAGE:2384491 3' similar to TR:P87890 P87890 POL PROTEIN :	Homo sapiens chromosome 21 segment HS21C027	QV1-BT0631-040900-357-f02 BT0631 Homo sapiens cDNA	7e34c03.x1 NCI_CGAP_Lu24 Homo sepiens cDNA done IMAGE:3284356 3' similar to WP:R151.6 CE00828 :	Homo sapiens X linked anhidroitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions
gle Exon Pro	Top Hit Database Source	ΗN	EST_HUMAN	FZ	EST_HUMAN	٦	NT	70.1 EST_HUMAN	Ę	NT	Ę	NT	TN	LX.	LN L	LN.	Ë	Z	EST HUMAN	1 EST_HUMAN	ᄓ	LN	Ę	TA.	EST HUMAN	뉟	EST_HUMAN	EST_HUMAN	Ņ
uis	Top Hit Acession No.	38 AF190501.1	38 AV726988.1	3.1	38 H55641.1	38 S74906.1	11418248 NT	38 AA401570.1	4885288	7881969 NT	38 AF270831.1	38 AL163203.2		8922543 NT	7305360 NT	7305360 NT		-38 11422250 NT			7662109	38 AL163284.2	4502312 NT	4758229 NT	8.0E-39 AI823404.1		6.0E-39 BF331829.1	6.0E-39 BE670394,1	5.0E-39 AF003528.1
	Most Similar (Top) Hit BLAST E Value	2.0E-38	2.05-38	2.0E-38	2.0E-38	2.0E-38	2.0E-38	1.0E-38	1.0E-38	1.0E-38		1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.0E-38	1.05-38	1.0E-38	1.0E-38	1.0E-38	8.0E-39	8.0E-39	8.0E-39	7.0E-39	6.0E-39	6.0E-39	5.0E-39
	Expression Signal	3.52	3.05	2.08	6.45	1.43	3.76	2.55	2.63	1.11	2.34	1.23	1.23	ŀ	4.71	4.71	3.15	0.71	5.13	0.58	1.28	2.2	15.3	1.45	1.27	5.79	2.24	1.86	1.57
	ORF SEQ ID NO:	37558			31081				27450	27475	27960			29921	31677		အ၀ငေ	34898	35150		37503			26796		27650	36639		26412
	Exon SEQ ID NO:	24235		24518	24705	_	25031	13829	14728	14747	15216	17029	17029	17293	18719		19987	21740	21977	22948	24187	25140	12882	14121	14560	14819	23400	24979	13750
	Probe SEQ ID NO:	11638	11971	11973	12260	12323	12777	1071	1992	2012	2499	4290	4290	4558	5937	5937	7304	9051	9310	10301	11588	12118	53	1373	1821	2087	10711	12696	987

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Top Hit Detabase Source EXPLINESSOUTH DITAIN Top Hit Detabase Source Colon HPLRB7 Home sapiens cDNA clone IMAGE:2374063 3' amiliar to TR:C15408 (15408 NEUTRAL) PROTEASE LARGE SUBEINIT conclains LTR7.11 LTR7 repetitive element; 12281 HUMAN Home sapiens chromosome 21 segment H52/C010 NT Home sapiens Planch for prostacyclin synthase, exen 2 NT Home sapiens chromosome 21 segment H52/C010 NT Home sapiens Real CTR repetitive element; 17 Home sapiens CDNA clone IMAGE:1020438 3' shrillar to contains CTR7.11 LTR7 repetitive element; 17 NT HOME sapiens Real CTR2 responsed to the Sapiens CDNA clone CR12-1 (17 NT CR14 CTR2 repetitive element); 17 NT HOME sapiens Real CTR2 responsed to the Sapiens CDNA clone CR12-1 (17 NT CR14 CTR2 repetitive element); 17 NT HOME sapiens Real CTR2 responsed protein 1 (RANGAP1), mRNA CH04 CH06 CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone CR12-1 (17 Regional genomic DNA specific CDNA library Home sapiens CDNA clone MAGE:160098 3' shrillar to SW1-3TR5_RAT (18 LCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4427 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TYPE 5, SMALL INTESTINE; 1 R4447 CLUCCSE TRANSPORTER TY	Top Hit Databese Source Source Source Source Source INT INT INT INT INT INT INT INT INT INT	_ _ % _ ` _		Expression Signal 8.62 2.04 6.78 0.97 0.97 0.97 0.95 0.95 0.97 0.95 0.95 0.95 0.95 0.95 0.95 0.95 0.95	R _O		Probe SEQ ID NO: 2884 12410 637 46 46 46 46 46 46 46 46 46 46 46 46 46
neutra incregious pudatve rout report for mentral incress, compress constitution of the second amage of 1.51 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:1630196 3'	EST_HUMAN	2.0E-39 AH0/8//8.1 2.0E-39 AA984531.1	1				8209 8394
Rattus norvegicus putative four repeat ion channel mRNA, complete cds	NT	AF078779.1			L	L	8209
Homo sapiens chromosome 21 segment noctions	-N	AL163202.2	ı				7431
Homo sapiens chromosome 21 segment Hox10002	F	AL163202.2	Ì				7431
ZNO602_r1 Stratagene hN I neuron (#43/233) Homo sapiens CUINA cione ilwa occione o	EST_HUMAN	AA080867.1					7269
ng86f03.s1 NCI_CGAP_Pr6 Homo sapiens cDNA clone IMAGE:941693	EST HUMAN	AA508880.11					5403
RC4-FN0037-290700-011-a10 FN0037 Homo saplens cDNA	EST HUMAN	BF370207.1					4370
Homo sapiens chromosome 21 segment HS21C048	. 1	AL163248.2				l _	2634
THR repetitive element;	EST_HUMAN	AA720574.1					1966
nw21g02.s1 NCI_CGAP_GCB0 Homo sapiens cDNA clone IMAGE:1241138 3' similar to contains THR.t3						L	
PM0-B10340-211299-003-402 B10340 Homo saplens cDNA	EST_HUMAN_	AW372318.1		11.91			1520
Homo sapiens homogentisate 1,2-dioxygenase gene, complete cds	TN.	AF000573.1		4.2		ľ	1009
promma-7 D01.r bytumor Homo saplens cDNA 5	EST HUMAN	AI525119.1		14.08		l	892
601301607F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3636289 5	EST_HUMAN	BE409203.1		5.8		13646	877
yp51c06.s1 Scares retina N2b4HR Homo saptens cDNA clone IMAGE:180954 3	EST_HUMAN	H37903.1		5.82		<u> </u>	12008
P43427 GLUCOSE TRANSPORTER TYPE 5, SMALL INTESTINE;	EST_HUMAN	A1084557.1					11963
0x83a10.s1 Soures NhHMPu S1 Homo saplens cDNA clone IMAGE:1660986 3' similar to SW:GTR5_RAT	i						3
OXXXXXIV.ST SOZIES_NITRIMITY_ST FORD SEPTENTS CLIVA CATE INVACE: 1000500 5 SUITED TO SOV.CO. N. J. N.	EST HUMAN	A1084557.1					11083
firmfor16 Regional genomic DNA specific cDNA library Homo sapiens cDNA cione CK12-1	EST HUMAN	AA631949.1					48
frific16 Regional genomic DNA specific cDNA library Homo sapiens cDNA clone CR12-1	EST_HUMAN	AA631949.1	3.0E-39				48
Ifmfc16 Regional genomic DNA specific cDNA library Homo sapiens cDNA clone CR12-1	EST_HUMAN	AA631949.1	8-30.E			١.	48
QV0-FN0063-260600-278-c06 FN0063 Homo sapiens cDNA			4.0E-39	2.71		L	12534
Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	. TN	11418177	4.0E-39	4.47		L	12427
Homo sapiens DNA for prostacyclin synthase, exon 2	NT					L	9228
Homo sapiens DNA for prostacyclin synthase, exon 2	N						9228
OFR.b1 OFR repetitive element;	—′	3.1	4.0E-3	1.27			7974
ae92g04.s1 Stratagene schizo brain S11 Homo sapiens cDNA done IMAGE:1020438 3' strullar to contains							
Homo sapiens chromosome 21 segment HS21C010	N					1	3559
Chlorocebus æethlops mRNA for ribosomal protein S4X, complete cds	NT					L	537
Homo saplens hypothetical protein FLJ10803 (FLJ10803), mRNA	NT	11420289	5.0E-39				12410
at36b04.x1 Barstead colon HPLRB7 Homo saplens cDNA clone IMAGE:2374063 3' stimitar to 1 R:Q15408 Q15408 NEUTRAL PROTEASE LARGE SUBUNIT ;contains LTR7.t1 LTR7 repetitive etement;	EST_HUMAN	AI750154.1	5.0E-3(2984
artishot vt Baretsad colon HPI RB7 Homo sapiens cDNA cione IMAGE:2374083 3' similar to TR:Q15408							
Top Hit Descriptor	l op Hit Database Source	Top Hit Acession No	(Top) Hit BLAST E Value	Expression Signal			Probe SEQ ID NO:
Des Expressed III Dialii		1110					

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Single Exon Probes Expressed in Brain	Most Similar Top Hit Acession (Top) Ht Top Hit Database Ignal BLAST E No. Source	0.73 2.0E-39 A1886860.1 EST HUMAN IN35e03.x1 NCI CGAP Prz8 Homo sapiens cDNA clone IMAGE-2253052.3	2.0E-39 D86964.1 NT	1.0E-39 AJ006345.1 NT	NT	19 7657020 NT	0303 NT	0303 NT	1.0E-39 AW951995.1 EST_HUMAN	39 AW951995.1 EST HUMAN	020 NT	1.0E-39 11417342 NT (TM) and short cytoplasmic domain, seven thrombospondin repeats (type 1 and type 1-like), transmembrane domain (semaphorin) 54 (SEMA5A), mRNA	1.0E-39 11417342 NT (TM) and short cytoplasmic domain. (semenborin) 5A (SEMA5A). mRNA		4 OE-30 A 12784704	- LV	1.0E-39 11436736	TN	SWISSPROT	1.0E-39 U07000.1 NT	9.0E-40 6803210 NT	755145 NT	15.14 9.0E-40 4755145 NT Homo sapiens AE-binding protein 1 (AEBP1) mRNA	6.54 9.0E-40 4507512 NT MRNA mRNA	0.97 9.0E-40 4503764 NT Homo sapiens fragile X mental retardation 1 (FMR1) mRNA	۲N	0.84 8.0E-40 AA078165.1 EST_HUMAN 7H15A04 Chromosome 7 HeLa cDivA Library Homo saplens cDNA clone 7H15A04	8.0E-40 BE396541.1 EST_HUMAN	2.03 7.0E-40 U60325.1 NT Human DNA polymerase gamma mRNA, nuclear gene encoding mitochondrial protein, complete cds
}	<u></u>	2.0E-39 AIG	2.0E-39 D8¢	1.0E-39 AJC	1.0E-39 AJC	1.0E-39	1.0E-39	1.0E-39	1.0E-39 AW	1.0E-39 AW	1.0E-39	1.0E-39	1.0E-39	1 OF 20 TO	1 0E-30 A 12	1 OF 30 A 17	1.0E-39	1.0E-39 D78	1.0E-39 O46	1.0E-39 U07	9.0E-40	9.0E-40	9.0E-40	9.0E-40	9.0E-40	9.0E-40 ABC	8.0E-40 AA0	8.0E-40 BE3	7.0E-40 U600
	Expression (*) Signal B	0.73	2.97	3.71	3.71	4.24	2.0	0.7	2.5	2.5	8.86	1.02	1.02	70	78.7	A 84	1.57	L					15.14	6.54	76.0	3.99	0.84	3.35	2.03
	ORF SEQ ID NO:		37365	26936	26837	26952	29467	29468	28980	29981	30021	30735	30736	31248	31287	31288	-	33025	34297		26957	26629	26630	26865	29155	29343	28449		33390
	Exan SEQ ID NO:	22179	24058	14249	14249	14268	16841	16841	17347	17347	17388	18079	18079	18330	18375	18375	19561	19948	21154	24761	13325	13983	13963	14179	16517	17878	15802	16853	20282
	Probe SEQ ID NO:	9628	11409	1503	1503	1521	4098	4098	4612	4612	4654 42	5274	5274	5542	5578	8578	6727	7264	8462	12357	£42	1213	1213	1432	3765	3956	3036	3903	7616

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					5	Sid Land	ייים יייים
Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7616	20282	33391	2.03	7.0E-4	to U60325.1	· FN	Human DNA polymerase gamma mRNA, nuclear gene encoding mitochondrial protein, complete cds
10813		36732	2.27	7.0E-40	10 AL163246.2	NT	Homo saplens chromosome 21 segment HS210046
2730	15437	28174	8.41	6.0E-40	6.0E-40 AA361275.1	EST_HUMAN	EST70527 T-cell lymphoma Homo sapiens cDNA 5' end similar to similar to zinc finger protein family
2730	15437	28175	8.41	6.0E-40		EST_HUMAN	EST70627 T-cell lymphoma Homo saplens cDNA 5' end similar to similar to zino finger protein family
5849		Ŀ		~30.9	BE504766.1	EST_HUMAN	hz40g01 x1 NCI_CGAP_GC8 Hamo sepiens cDNA clone IMAGE:3210480 3'
6055	L		1.11	6.0E~	7661999 NT	NT	Homo sapiens KIAA0211 gene product (KIAA0211), mRNA
6836	19498	32522	3.56	6.0E-40	11439783 NT	NT.	Homo sapiens fatty acid desaturase 1 (FADS1), mRNA
6836	19498	32623			11439783 NT	Ŋ	Homo sapiens fatty acid desaturase 1 (FADS1), mRNA
9877	L				6.0E-40 AV653028.1	EST_HUMAN	AV653028 GLC Homo sapiens cDNA clone GLCDGF04 3'
9877	L	35723			6.0E-40 AV653028.1	EST_HUMAN	AV653028 GLC Homo sapiens cDNA clone GLCDGF04 3'
99,	4,607		1 70	7 0 7	A SERBONE 1	NAMIH TAR	#91b01.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2248873 3' similar to TR:O73505 O73505 POI_PROTEIN_:
300	1			1			Home ceniess X-linked anhidmitic extraternal dissulasta protein cana (EDA), exon 2 and flanking repeat
2101	14832		2.27	4.0E-4	10 AF003528.1	뉟	וויסווס פקומוס אל יווישט פווישט פער אייני אייני פער אייני אייני אייני אייני אייני אייני אייני אייני אייני אייני
4356		29729			7662117 NT	N	Homo sapiens KIAA0433 protein (KIAA0433), mRNA
7786	<u> </u>	L	0.5		4.0E-40 AU127831.1	EST_HUMAN	AU127831 NT2RP2 Homo sapiens cDNA clone NT2RP2002172 5'
7890					4.0E-40 AA742809.1	EST_HUMAN	nv34e10.r1 NCI_CGAP_Br4 Homo saplens cDNA clone IMAGE:1222122
8953	L	L	5.17		BE009416.1	EST_HUMAN	PM0-BN0167-070500-002-h12 BN0167 Homo saplens cDNA
8953	L	L			4.0E-40 BE009416.1	EST_HUMAN	PM0-BN0167-070500-002-h12 BN0167 Homo saplens cDNA
10616	L				AW841585.1	EST_HUMAN	RC1-CN0017-120200-012-e04 CN0017 Homo sapiens cDNA
4111	16854		1.02	Ĺ	3.0E-40 A1925949.1	EST_HUMAN	wh12f07.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2380549 3'
	_						Homo sapiens sema domain, seven thrombospondin repeats (type 1 and type 1-like), transmembrane domain
6543	19308	32313	7.02	3.0E~	•	ΝΤ	(TM) and short cytoplasmic domain, (semaphorin) 54 (SEMASA), mKNA
828					5454167 NT	N	Homo sapiens HBV associated factor (XAP4) mRNA
8888	L		1.25	3.0E-4	ID AF078779.1	N	Rattus norvegicus putative four repeat ion channel mRNA, complete cds
9111	<u>L</u> .			3.0E-	to AF078779.1	N	Rattus norvegicus putative four repeat ion channel mRNA, complete cds
11232	L	37182	8.36	3.0E-	6005813 NT	NT	Homo saplens serine threonine protein kinase (NDR), mRNA
44583	24482	27775	2.23	6. H	00 AW118799 1	FST HUMAN	xd96h02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2605491 3' similar to TR:Q15804 O15804 SIMILAR TO ENV OF TYPE A AND TYPE B RETROVIRUSES AND TO CLASS II HERVS;
				200	A1222038 4	EST HIMAN	ans 2 hours testis NHT Homo saniens cDNA clone IMAGE:1838847 3
31/	13120		3.0	Z.0E-	#0 MIZZOWO. 1	TOWOL TO TO	

Page 280 of 536 Table 4 Single Exon Probes Expressed in Brain

Γ		_	T	_	÷	T _e	7	_	_	Т	Т	Т	1	_	_	_	T	_	_	_	т_	· T	_	· ·	_				T-
Single Exon Probes Expressed in Brain	Top Hit Descriptor	xr24e10.x1 NCI_CGAP_UM Homo septens cDNA clone IMAGE.2781088 3' similar to SW:RS5_MOUSE P97461 40S RIBOSOMAL PROTEIN S5.;	AV731601 HTF Homo saplens cDNA clone HTFAZE05 5'	Homo saplens proteasome (prosome, macropain) subunit, alpha type, 7 (PSMA7) mRNA, and translated products	Homo sepiens proteasome (prosome, macropain) subunit, alpha type, 7 (PSMA7) mRNA, and translated products	w80a11.x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:2814716 3' similar to TR:Q91929 Q91929 ZNC FINGER PROTEIN.	Homo saplens adently cyclase-associated protein 2 (CAP2) mRNA	601121567F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3345784 6"	Homo sapiens adenylyl cyolase-associated protein 2 (CAP2) mRNA	Homo saplens chromosome 21 segment HS21C080	Homo saplens chromosome 21 segment HS21C080	nc09a09.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1007808	601460375F1 NIH_MGC_68 Homo sepiens cDNA clane IMAGE:3863803 5'	bb79a10.71 NIH_MGC_10 Homo saplens cDNA clone IMAGE:3048570 5' similar to TR:Q9Z188 Q9Z168 SYNTAXIN 17.;	602068604F1 NIH_MGC_58 Homo saplens cDNA clone IMAGE:4067736 5'	602068604F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4067736 5	Homo septens sorting nextn 3 (SNX3) mRNA	Homo saplens zinc finger protein 200 (ZNF200) mRNA, and translated products	zh79f11.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:418317 3'	zh79f11.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:418317.3'	nj42f04.s1 NCI_CGAP_AA1 Homo sapiens cDNA clone IMAGE:995167 3'	nj42f04.s1 NCI_CGAP_AA1 Homo saplens cDNA clone IMAGE:995187 3'	POL POLYPROTEIN (CONTAINS: PROTEASE ; REVERSE TRANSCRIPTASE ; RIBONUCLEASE H 1	AU148345 NT2RM4 Homo saplens cDNA clone NT2RM4002122 3'	np09h03.s1 NCI_CGAP_Pr3 Homo sapiens cDNA clone IMAGE:1115861 similar to TR:G1138406	G1138406 KIAA0173 PROTEIN.;	np09h03.s1 NCL_CGAP_Pr3 Homo sepiene oDNA clone IMAGE:1115861 similar to TR:G1136406 G1136406 KIAA0173 PROTEIN.:	MR2-CT0222-211089-002-610 CT0222 Homo sapiens dDNA	Homo saplens chromosome 21 segment HS21C003
JIO EXON Proi	Top Hit Database Source	EST_HUMAN	EST_HUMAN	TN	۲	EST HUMAN	FZ	EST_HUMAN	N	LN	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	N	NT		EST_HUMAN		EST_HUMAN	SWISSPROT	EST_HUMAN		EST_HUMAN	EST HUMAN	Т	NT
מוט	Top Hit Acession No.	2.0E-40 AW303868.1	0 AV731601.1	4506188 NT	450B188 NT	Al968562.1	5453592 NT	BE275932.1	5453592 NT	AL163280.2	AL163280.2	AA225989.1	BF036881.1	BE018348.1	BF541030.1	BF541030.1	4507142 NT	4508012 NT	W92708.1	W92708.1	AA573201.1	AA573201.1	P26808	AU148345.1		AA614255.1	AA614255.1		
	Most Similar (Top) Hit BLAST E Value	2.0E-40	2.0E-40	2.0E-40	2.0E-40	2.0E-40	2.0E-40	-	2.0E-40	2.0E-40 /	2.0E-40 /		1.0E-40 E	1.0E-40 E	1.0E-40 E	1.0E-40 E	1.0E-40	1.0E-40		1.0E-40 V	1.0E-40 A	1.0E-40 A	1.0E-40 F	1.0E-40		1.0E-40 A	1.0E-40		8.0E-41 A
	Expression Signal	1.61	0.92	1.58	1.58	1.21	2.48	1.44	4.28	1.68	1.68	1.78	0.93	\$.	1.18	1.18	1.27	4.52	0.75	0.75	1.77	1.77	0.89	8.34		1.89	1.89	10.09	1.62
	ORF SEQ ID NO:			27376	27376	27622	27630		28529	30197	30198		28083		28185	28186		29934	31907	31908	32727	32728	32886	36758		37612	37613		33643
	Exon SEQ ID NO:	13549	14557	14663	14663	14796	14895	15404	15888	17573	17573	13632	15339	15401	15447	15447	16053	17306	18938	18938	19880	19680	19820	23516		24289	24289	25274	20517
	Probe SEQ ID NO:	111	1818	1927	1927	2084	2166	2692	3123	4843	4843	863	2627	2692	2741	2741	3292	4571	6161	6161	6987	6987	7133	10834		11694	11694	12376	7822

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						200	
Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression . Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
808	15553	L	1.24	7.0E-41	7.0E-41 A1934364.1	EST_HUMAN	wp04h04.xrl NCI_CGAP_Kid11 Hamo sapiens cDNA clone IMAGE:2463895 3'
808		26247	1.24	7.0E-41	7.0E-41 AI934364.1	EST_HUMAN	wp04h04.x1 NCI_CGAP_Kid11 Hamo sapiens cDNA clane IMAGE:2463895 31
4609	17344	29976	0.92	7.0E-41	1 BE389592.1	EST_HUMAN	601282077F1 NIH_MGC_44 Homo sepiens cDNA clone IMAGE:3603955 5'
4609	17344	29977	0.92	7.0E-41	7.0E-41 BE389592.1	EST_HUMAN	601282077F1 NIH_MGC_44 Hamo sapiens cDNA clone IMAGE:3603955 5'
5183	17991	30507	1.2	7.0E-41		TN	Homo sapiens hypothetical protein FLJ13188 (FLJ13188), mRNA
5918	18703	31656	3.49	7.0E-41	11419208 NT	NT	Homo sapiens a disintegrin and metalloproteinase domain 22 (ADAW22), mRNA
6260	19034	. 1	0.61	7.0E-41	11433010 NT	LZ	Homo sapiens IQ motif containing GTPase activating protein 1 (IQGAP1), mRNA
6895	17971	30528	89'0	7.0E-41	7.0E-41 U72335.1	F	Human platelet activating factor acety/hydrolase, brain isoform, 45 kDa subunit (LIS1) gene, exons 3 and 4
11411	ı			7.0E-41	4758445 NT	NT	Homo saplens guanine nucleotide binding protein 10 (GNG10) mRNA
	<u> </u>			77 20 2	. 10000001	j.	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, excns 7-49, and partial cds, alternatively
1831		37552		7.0E-41	AF 2233	z !	spineso
12782	25271		4.35	7.0E-41	11417972 NT	L _Z	Homo septiens pescadillo (zebrarish) nomalog 1, contaming BRC1 algorithms.
274	13081	25724		6.0E-41	AB03716	LN	Homo sapiens DSCR5b mRNA, complete cds
2104	14835		2.04	6.0E-41	7657042 NT	NT	Homo sapiens Down syndrome candidate region 1 (DSCR1), mRNA
4433	17169	29797	16.0	6.0E-41	6.0E-41 BE567816.1	EST_HUMAN	601340485F1 NIH MGC_53 Hamo sapiens cDNA clone IMAGE:3682677 5
7871	20566		1.44	6.0E-41	6.0E-41 BF513783.1	EST_HUMAN	UFH-BW1-amp-b-03-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3070421 3'
1795	14535	27244	1.11	5.0E-41	T62628.1	EST_HUMAN	yc03e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:79626 3'
4087	16830		1.07	5.0E-41	4885636 NT	LΝ	Homo sapiens target of myb1 (chicken) homolog (TOM1), mRNA
6452	19220		2.29	5.0E-41	41 BE067042.1	EST_HUMAN	PM4-BT0341-251199-002-F11 BT0341 Homo sapiens cDNA
382	13169		2.42	4.0E-41	41 BE156318.1	EST_HUMAN	QV0-HT0367-150200-114-g09 HT0367 Homo saplens cDNA
1076	13834	26492	1.26	4.0E-41	41 AU119344.1	EST_HUMAN	AU119344 HEMBA1 Homo sapiens cDNA clone HEMBA1005583 5'
1388	14135	26810	15.51	4.0E-41	41 AI027117.1	EST_HUMAN	ow45e06.s1 Soares_parathyrold_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1649794 3' similar to TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE. ;contains LTR5.b1 LTR5 repetitive element ;
	ŧ						ow45e06.s1 Soares_parathyraid_tumor_NbHPA Homo saptens cDNA clone IMAGE:1849794 3' similar to
1388	14135	26811	15.51	4.0E-41		EST_HUMAN	TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE.; contains LTR5.b1 LTR5 repetitive element;
1403				4.0E-41	41 AB008681.1	NT	Homo saplens gene for activin receptor type IIB, complete cds
1632				4.0E-41	41 AI500406.1	EST HUMAN	tm96c04.x1 NCI_CGAP_Brn25 Homo saplens cDNA clone IMAGE:2165958 3' similar to contains OFR.b1 OFR repetitive element ;
2891	1_	L		4.0E-41	41 AJ229041.1	ΓN	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3
2891	<u>L</u>			4.0E-41	41 AJ229041.1	NT	Hamo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3
4124	li	29493		4.0E-41	41 X92685.1	NT	H. sapiens DNase I hypersensitive site (HSS-3) enhancer element

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- 1		Т	_	1	_	т	_	_	_	1	_	_	τ	_	_	_	_	_	_	_	_	_	, –	_	_		_	_	_		_	_	
Single Exon Probes Expressed in Brain	Top Hit Descriptor	AV758295 BM Homo saplens cDNA clone BMFBHC06 5'	601888096F1 NIH MGC 17 Home saplens cDNA clone IMAGE:4122119 5'	AV710480 Cu Homo sapiens cDNA clone CuAACCO7 5'	AV708431 ADC Homo sapiens cDNA clone ADCARE02 5"	601508315F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3910059 6'	Homo sapiens PAD-H19 mRNA for peptidy/arginine deminase type II, complete cds	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	H.saplens mRNA for putative p84 CLCP protein	Homo saplens mRNA for KIAA1387 protein, partial cds	EST64683 Jurkat T-cells VI Homo sapiens cDNA 5' end	Homo saplens 959 kb contig between AML1 and CBR1 on chromosome 21a22; seament 1/3	af17/10.s1 Soares, testis, NHT Homo saplens cDNA clone IMAGE:1031947.3	601762940F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:4026081 5'	Human ribosomal protein L23a mRNA, complete cds	EST35818 Embryo, 8 week I Homo saplens cDNA 5' end	Human mRNA for KIAA0207 gene, complete cds	G.gorilla DNA for ZNF80 gene homolog	Human ribosomal protein L23a mRNA, complete ods	208b04.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:785839 5	Homo saplens chromosome 21 segment HS21C067	Homo sapiens chromosome 21 segment HS21C067	xm4708 x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2687363 3' similar to TR:O70343 O70343	Homo saalens inferrin beta 8 (ITGRs) mRNA	Homo sabiens homolog of Nedd5 (hNedd5) mRNA complete cits	Human B-cell specific transcription factor (RSAP) mRNA complete eds	Human B-cell specific transcription factor (BSAP) mRNA, complete cds	EST31723 Embryo, 12 week I Hamo saplens cDNA 5' end	ZINC FINGER PROTEIN 135	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	EST84555 Colon adenocarcinoma IV Homo sapiens cDNA 5' end	601445647F1 NIH_MGC_65 Homo saplens cDNA done IMAGE:3849803 5'
gie Exon Pro	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	NT	LN	LN	Z,	EST_HUMAN	Ę	EST HUMAN	EST_HUMAN	Z	EST_HUMAN	F	Ę	LN LN	EST_HUMAN	IN	NT	TOT LIMAN	NAMOL 100	Į.	NT	Z-	EST_HUMAN	SWISSPROT	F	NT	П	EST_HUMAN
uio	Top Hit Acession No.	-41 AV758295.1	-41 BF304683.1	AV710480.1	4.0E-41 AV708431.1	-41 BE887118.1	41 AB030176.1	41 AB026898.1		15.	41 AA356168.1		41 AA609768.1		41 U43701.1	41 AA331940.1	41 D86962.1	41 X89631.1	41 U43701.1			41 AL163267.2		41 AV 230347.1 E3	41 AF038404.1	Γ		41 AA328265.1	41 P52742	11417118	11417118 NT		41 BE869735.1
	Most Similar (Top) Hit BLAST E Value	4.0E-41	4.0E-41	4.0E-41	4.0E-41	4.0E-41	3.0E-41	3.0E-41	3.0E-41	3.0E-41	3.0E-41	3.0E-41	3.0E-41	3.0E-41	2.0E-41	2.0E-41	20E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41	77 30 6	2.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41			1.0E-41
	Expression Signal	1.41	7.24	11.07	1.63	1.69	2.68	2.45	7.78	1.59	0.71	1.28	1.52	1.48	5.17	1.84	1.54	3.34	4.65	1.41	1.17	1.17	0	0.76	8.08	1.45	1.45	1.12	1.61	0.52	0.52	2.76	1.05
	ORF SEQ ID NO:		35430				26358	29667	30908	32043	32916	37647			26987	27399	27684	27730	26987	28731	29942	29943	30475	32300	33348	33786	33787	33818	34710	35155	35156	37379	28616
	Exon SEQ ID NO:	19184	22246	24266			13694	17040	18204	19061	19846	24323	24485	24825	14299	14686	14944	14990	14299	16081	17314	17314	17850	19296	20241	20662	20662	20690	21565	21984	21984	24071	15964
	Probe SEQ ID NO:	6416	9593	11671	12646	12727	927	4301	5404	6288	7159	11730	11924	12456	1817	1951	2216	2264	2831	3321	4579	4679	5141	6530	7672	7967	7967	7995	8874	9317	9317	11468	3201

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						2.6	
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEO ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
3201	15964	28617	1.05	1.0E-41	1 BE869735.1	EST_HUMAN	601445647F1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3849803 5'
4529	17264	29897	14.08	1.0E-41	F678468	INT	Mus musculus tubulin alpha 6 (Tuba6), mRNA
6749	L	30582	99'0	1,0E-41	1 H99079.1	EST_HUMAN	yx18b03.s1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:262061 3'
9318	I		1.69	1.0E-41	11 A1217868.1	EST_HUMAN	q775c10.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1755858 3'
11111	┖			1.0E-41	1.0E-41 AW847812.1	EST_HUMAN	IL3-CT0213-190200-040-F09 CT0213 Homo saplens cDNA
12054	乚		2.81	1.0E-41	11526291		Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA
8418	_		1.14	9.0E-42	12 BE179191.1	EST_HUMAN	RC0-HT0613-210300-032-g01 HT0613 Homo sapiens cDNA
9072	21761	34922	3.49	9.0E-42	11560151 NT	TN	Homo saplens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA
9072		L	3.49	9.0E-42	11560151 NT	L	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA
450	13236	25875	7.71	8.0E-42	8.0E-42 AF003530.1	NT	Homo saplens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions
8				0 0 0 4 2	0 0 0 10 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	FIX	Hamo septiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete arts)
7017	4655	/00/7	0.92	0.05-42	AD020090.1	2	POSTIPINA CONTRACTOR THAT HOLD STATE HAVE THAT CONTRACTOR SIMILAR TO CARAGOA CARAGOA
12093	25277		4.4	8.0E-42	2 AA493896.1	EST_HUMAN	http://dx.sh.nci_coatingrinders edited content of the second similar to increase the second
	L				١		xc97a04.x1 NCI_CGAP_Brn35 Homo sapiens cDNA clone IMAGE:2592174 3' shnilar to contains OFR.t2
12111	25154		1.56	8.0E-42	2 AW088062.1	EST_HUMAN	OFR repetitive element;
911	13678		2.58	7.0E-42	12 AL163285.2	IN	Homo sapiens chranosome 21 segment HS21C085
9143	21874	35039		7.0E-42	12 AI204358.1	EST_HUMAN	qf58g12.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1754278 3'
11126	L_	l	1.3	7.0E-42	12 AA569592.1	EST_HUMAN	nf23g07,s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:914652
11126	3 23795		1.3	7.0E-42	7.0E-42 AA569592.1	EST_HUMAN	nf23g07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:914652
1848	L		6	6.0E-42	42 AF012872.1	NT	Homo sapiens phosphatidylinositol 4-kinase 230 (pl4K230) mRNA, complete cds
1848	ı	27300		6.0E-42	42 AF012872.1	NT	Homo sapiens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds
						100	xp29f08.x1 NCI_CGAP_HN10 Homo sapiens cDNA clone IMAGE:2741799 3' similar to contains L1.t1 L1
Ì		١	2,33	0.05-42	12 AW 230000.1	ESI _ IDOINGN	II promise and N. Ver VIAAGEZ meetin partial cale
5381		30871	1.63	6.0E-	١	N .	THAT SEPTICAL STANKS AND AND AND AND AND AND AND AND AND AND
5630	_		1.45			LN-	Homo sapiens mKNA for KIAA 1067 protein, partial cas
132	12947		7.53	5.0E-42	42 AJ271735.1	NT	Homo saplens Xq pseudoautosonal region; segment 1/2
428		25859	1.41	5.0E-42	5.0E-42 BE217913.1	EST_HUMAN	hv31e11.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3175052 3'
474	_		2.57	5.0E-42	5730038 NT	NT	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA
475	L		2.74	5.0E-42	F730038 NT	NT	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA
	}					<u>.</u>	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman
6587	19350	32363	1.04	5.0E-42	11433063 N	Z	Synaromej (OBESA), mrvva
6587	19350	32364	1.04	5.0E-42	11433063 NT	Ę	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A), mRNA
	ı				١		

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens myotubularin related protein 3 (MTMR3), mRNA	Homo sapiens mulitfunctional calcium/calmodulin-dependent protein kinase II delta2 isoform mRNA, complete cds	Homo saplens reelin (RELN) mRNA	Homo sapiens mRNA for KIAA1294 protein, partial cds	Homo saplens hypothetical protein FLJ20163 (FLJ20163), mRNA	Homo sapiens MHC class 1 region	Homo saplans MHC class 1 region	Homo sapiens ribonuclease III (RN3) mRNA, complete cds	H.sapiens PROS-27 mRNA	Homo saplens SNARE protein kinase SNAK mRNA, complete cds	Homo sapians regulatory factor X, 4 (influences HLA class II expression) (RFX4) mRNA	Homo saplens zinc finger protein 177 (ZNF177) mRNA	RC1-ST0278-040400-018-h11 ST0278 Homo sapiens cDNA	RC1-ST0278-040400-018-h11 ST0278 Homo sapiens cDNA	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	RC0-TN0079-110900-024-g07 TN0079 Homo saplens cDNA	RC3-NN0070-270400-011-h10 NN0070 Homo sapiens cDNA	2819293.3prime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2819293.3'	EST367438 MAGE resequences, MAGC Homo sapiens cDNA	EST367438 MAGE resequences, MAGC Homo sapiens cDNA	ow83d05.x1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1653417 3'	601061284F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3447620 5'	RIBONUCLEASE K3 (RNASE K3)	RIBONUCI EASE K3 (RNASE K3)	Homo sapiens chromosome 21 segment HS21C046	Human endogenous retrovirus pHE.1 (ERV9)	UI-H-BI1-afh-e-04-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:27218713'	Homo sapiens partial C9 gene for complement component C9, exon 1	Homo saplens partial C9 gene for complement component C9, exon 1	Homo sepiens NADH-ubiquinone oxidoreductase AGGG subunit precursor homolog mRNA, nuclear gene	encoding miliconoridatel protein, complete cds	raans sapens NAUT-ubiquinone oxidoreductase AGGG subunit precursor homolog mRNA, nuclear gene encoding mitochondrial protein, complete cds
gle Exon Prob	Top Hit Database Source	LN	LN.		Z	FZ	NT				본			EST_HUMAN		EST_HUMAN	EST_HUMAN		EST_HUMAN			EST_HUMAN		SWISSPROT	/ISSPROT			EST_HUMAN		INT		2	Į.
Sing	Top Hit Acession No.	11417957 NT	-42 AF071569.1	26977	5.0E-42 AB037715.1	8923162 NT		-42 AF055066.1	.1	-42 X59417.1	-42 AF246219.1	4506496 NT	4508008 NT	-42 AW818630.1	1		2.0E-42 BF376834.1		-42 AW 250059.1		1	2.0E-42 AI052586.1	19.1			16.2			1.0E-42 AJ251818.1			1.0E-42 Ar 06/166.1	42 AF067166.1
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Page 285 of 536 Table 4 Single Exon Probes Expressed in Brain

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	Expression	4.55	1.52	1.17	0.72	0.46	0.47	0.47	4.44	2.31	1.21	5.29	1.84	5.0	1.02	0.7	2.32	5.21	5.21	1.23	1.68	3.05	4.59	2.07	1.31	1.04
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	Probe SEQ ID NO:	490	2850	6213	6802	8778	9248	9248	9264	10297	10335	10683	1089	952	6178	6276	7030	8077	8077	10213	11275	12030	1191	1690	3558	4258

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Sin	Top Hit Acession No.	11418322 NT	8.0E-44 AI222985.1			1423497		Y10498.2	8.0E-44 L29139.1	327389	11418086 NT	11418099 NT	-44 R06035.1	5031886	-44 AF048729.1	Γ	Τ		ľ	5646	7.0E-44 AU159839.1	Γ			5.0E-44 AJ289880.1 N		T.	ľ		l		T	-
	Most Similar (Top) Hit BLAST E Value	9.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	8.0E-44	7.0E-44	7.0E-44	7.0E-44 /	7.0E-44 /	7.0E-44/	7.0E-44 /	7.0E-44	7.0E-44	7.0E-44 A	6.0E-44 Z20948.1	6.0E-44	5.0E-44 A	5.0E-44 A	4 4 4	5.0E-44 AISB8523.1	4 0F-44 A	4 0F 44 B	4 OF 44 1	4 0F-44 B	4.0E-44 U	3.0E-44 A
	Expression Signal	3.41	6.23	6.23	2.67	0.47	0.47	3.78	1.38	4.09	1.38	2.55	0.69	1.06	2.58	2.58	2.54	1.12	1.12	1.01	2.28	0.94	1.78	3.3	2.72	90	7.34	3.75	88 0	0.78	0.51	5.38	5.77
	ORF SEQ ID NO:	31032	26309	26310	34288	36097	36098	37084	37606	31073	30976	30808		27696	28381	28382	29231	29581	29582	30476	33908	31756	37702		1	33607	10000	28816	33131	33995		37153	28500
	Exon SEQ ID NO:	24835					22884	LI				25186	13423	14956	15732	15732		16958		17860	20779	18793	24372	13102	13124	20483	22038	16167	20050	20863	21469	23868	15859
	SEQ ID NO:	12469	870	870	8437	10238	10236	11117	11688	12207	12248	12589	4	2228	2968	2968	3843	4217	4217	5142	8085	6012	11781	987	323	7788	9284	3409	7370	8169	8777	11202	3094

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יייים ביייים או הספסה ביייים בייים ביייים יייים ביייים ביייים בייים ייים בייים בייים בייים בייים ביים בייים ייים בייים בייים בייים בייים בייים ביים בייים יים בייים יים בייים ביים ביים ביים בייים ביים ביים ביים	Top Hit Descriptor	EST42299 Endometrial tumor Homo sapiens cDNA 5' end similar to similar to alpha-1-entiproteinase F	Sus scrofa domestica submaxillary apomucin mRNA, complete cds	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1 (DDX1) mRNA	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1 (DDX1) mRNA	Homo sapiens transmembrane trafficking protein (TMP21), mRNA	Homo sapiens transmembrane trafficking protein (TMP21), mRNA	Homo sapiens RAB36 (RAB38) mRNA, complete cds	hw14g08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3182938 3' similar to SW:OXYB_HUMAN P22669 OXYSTEROL-BINDING PROTEIN ;	Homo sapiens tissue-type bone marrow zinc finger protein 4 mRNA, complete cds	Homo saplens adaptor-related protein complex 4, sigma 1 subunit (CLAPS4), mRNA	Homo sapiens DNA for amyloid precursor protein, complete cds	PM4-SN0016-120500-003-a04 SN0016 Homo sapiens cDNA	Homo saptens chemokine (C-C motif) receptor 9 (CCR9), mRNA	Homo sapiens general transcription factor 2-I (GTF2I) mRNA, atternatively spliced product, complete cds	Homo sapiens glutamate receptor, metabotropic 3 (GRM3), mRNA	Homo saplens glutamate receptor, metabotropic 3 (GRM3), mRNA	Homo sapiens vesicle transport-related protein (KIAA0917), mRNA	Homo saplens vesicle transport-related protein (KIAA0917), mRNA	801286914F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3613586 5	TCBAP1E2795 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project≃TCBA Homo sapiens cDNA clone TCBAP2795	Homo saplens cat eye syndrome chromosome region, candidate 1 (CECR1), mRNA	Homo sepiens Misshapen/NIK-related kinase (MINK), mRNA	Homo sapiens Misshapen/NIK-related kinase (MINK), mRNA	RC1-CT0249-030300-026-h12 CT0249 Homo sepiens cDNA	RC1-BN0039-110300-012-b01 BN0039 Homo sapiens cDNA	Homo saplens chromosome 21 segment HS21C103	zw53d02.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773763 5' similar to	contains THR.t3 THR repetitive element;	zw53d02.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone.IMAGE:773763 5' similar to contains THR.t3 THR to THR to the peritive element;	
מון ווסעדו פול	Top Hit Detabase Source	EST_HUMAN	NT	ΝT	LN	NT	NT	NT	EST_HUMAN	NT	N	Ę	EST_HUMAN	NT	Ä	NT	NT	NT	. TN	EST_HUMAN	EST_HUMAN	NT	N N	LN	EST_HUMAN	EST_HUMAN	NT		EST_HUMAN	EST HUMAN	
	Top Hit Acession No.	44 AA337234.1	44 AF005273.1	4826685 NT	4826685 NT	5803200 NT	5803200 NT	44 AF133588.1	44 BE465325.1	44 AF070651.1	5901933 NT	-44 D87675.1	44 AW864379.1	11449901 NT	44 AF038968.1	11419226 NT	11419226 NT	T706370 NT	7706370 NT	44 BE389058.1	44 BE244902.1	11526293 NT	7657334 NT	7657334 NT	44 AW853132.1	44 AW994803.1	44 AL163303.2	·	44 AA434554.1	44 AA434554.1	
	Most Similar (Top) Hit BLAST E Value	3.0E-44	3.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	2.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	-	1.0E-44	1.0E-44	
	Expression Signal	1.37	0.65	2.64	2.64	3.36	3.36	4.08	1.3	2.22	1.31	1.13	1.54	1.87	3.31	4.57	4.57	0.67	0.67	1.58	1.62	1.4	2.43	2.43	2.44	1.9	5.78		3.74	3.74	
	ORF SEQ ID NO:	29260	35269	26446	26447	26602	26603			Ì			29899		30562	33074	33075		34156	l.			L		25975				27687	27888	
	Exan SEQ ID NO:	16620	22097	13787	13787	13937	13937	14038	14095	14877	15327	16219	17266	18785	17927	•		21020	21020	21209	24456	25020	12880	12880	13347	13928	14314		14949	14949	2
	Probe SEQ ID NO:	3870	9419	1027	1027	1185	1185	1289	1347	2147	2616	3463	4631	8 9	6758	7313	7313	8327	8327	8517	11883	12760	હ	53	286	1175	1587		2221	2224	

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Top Hit Descriptor	zt88g11.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:7294765'	Homo saptens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM6 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, tiple LIM domein protein 6, and synaptophysin genes,	complete cds; and L-type calclum channel a>	aa01c09.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:811984 3'	Homo sapiens alpha satellite DNA, M1 monomer type	Homo sapiens alpha satellite DNA, M1 monomer type	EST378147 MAGE resequences, MAGJ Homo saplens cDNA	EST378147 MAGE resequences, MAGJ Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C009	qx88g07.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2009628 3'	AV714608 DCB Hamo saplens cDNA clone DCBBYE03 5'	Homo sapiens Sushi domain (SCR repeat) containing (BK65A6.2), mRNA	RC1-CT0198-150999-011-C08 CT0198 Homo saplens cDNA	RC1-CT0198-150999-011-C08 CT0198 Homo saplens cDNA	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sapiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sapiens mRNA for KIAA0995 protein, partial cds	Homo saplens TRK-fused gene (NOTE: non-standard symbol and name) (TFG) mRNA	Homo sapiens TRK-fused gene (NOTE: non-standard symbol and name) (TFG) mRNA	CM0-NN0005-130300-283-b09 NN0005 Homo saplens cDNA	EST90893 Synovial sarcoma Homo saplens cDNA 6' end	wb99c06.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2313802 3' similar to contains L1.t1 L1 respective element:	au83h07.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782809 3' similar to	SW R13A_HUMAN P40429 60S RIBOSOMAL PROTEIN L13A;	Homo saplens ADP-ribosylation factor GTPase activating protein 1 (ARFGAP1), mRNA	Homo saplens chromosome 21 segment HS21C003	CM4-CN0044-180200-515-f01 CN0044 Homo saplens cDNA	tg94f07.x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2116453 3' simitar to SW:PAX1_MOUSE P09084 PAIRED BOX PROTEIN PAX-1.;	z/22d03.st Sogres_testis_NHT Homo sapiens cDNA clone IMAGE:727877 3' similar to contains element TAR1 repetitive element;	Homo saplens MCP-1 gene and enhancer region
Top Hit Database Source	EST_HUMAN		NT	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	MT	EST HUMAN	EST_HUMAN	ΤN	NT	NT	NT	NT	EST_HUMAN	EST_HUMAN	FST HUMAN		EST_HUMAN	NT	LN	EST_HUMAN	EST HUMAN	EST HUMAN	Z.
Top Hit Acession No.	4 AA398099.1		4 AF196779.1	4 AA455869.1	4 AJ130755.1	4 AJ130755.1			4 AL163209.2	4 Al337183.1	4 AV714608.1	10092664 NT	4 AW846967.1	4 AW846967.1	8922391 NT	- 8922391 NT	5 AB023212.1	5174718 NT	6174718 NT	5 AW892763.1	8.0E-45 AA377985.1	5 A1875425 1		6 AW157570.1	11418213 NT	6.0E-45 AL163203.2	5.0E-45 BF333627.1	5.0E-45 AI523766.1	5 AA397781.1	5 Y18933.1
Most Similar (Top) Hit BLAST E Value	1.0E-44		1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44	1.0E-44 /	1.0E-44	1.0E-44	1.0E-44	1.0E-44	9.0E-45	9.0E-45	9.0E-45	8.0E-45	8.0E-45	8.0E-45	8.0E-45	8 OF 45		6.0E-45	6.0E-45	6.0E-45	5.0E-45	5.0E-45	5.0E-4	6.0E-4
Expression Signal	96'0		1.44	3.73	1.04	1.04	0.98	96'0	86'0	0.69	4.04	3.92	3.17	3.17	1.38	1.38	1.31	3.12	6.41	0.66	0.91	10,4		4.09	1.65	1.03	3.65	1.79		
ORF SEQ ID NO:	27746		28211		30385	30386	33988	33989	34380			37427	37496			28907	32323	L		Ŀ	33830						27453	28621		
Exon SEQ ID NO:	15590	L	15468	16485	17787	1		20857	21236	•	23616	24116	<u>L</u>	24182	i_	<u> </u>	18317	<u>l_</u>	L	19182	20701		1	16709	Ł	l	L	İ	<u> </u>	1 1
Probe SEQ ID NO:	2280		2763	3712	5048	5048	8163	8163	8544	8924	10936	11516	11583	11583	4639	4539	8552	2527	5023	6414	8008	1848	2	3960	12655	872	1995	3204	5425	6929

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Γ	- 1	-	. [Т	T	Т	Т	T	Т	Т	Т		T	T	Т	Т	Т	Т	Т	1	T	Т	Т	$\neg \neg$	T	T	T	٦		T	Т	Т	Т	٦
טווקם באכון רוססס באני ווי נימון	Top Hit Descriptor	Homo sapiens MCP-1 gene and enhancer region	Homo sapiens mRNA for inducible nitric codde synthase, complete cds	Homo sepiens mRNA for inducible nitric code synthase, complete cds	Homo sapiens zino finger protein 277 (ZNF277), mRNA	Homo sapiens zinc finger protein 277 (ZNF277), mRNA	Homo sapiens bone morphogenetic protein 5 (BMP5), mRNA	Homo sapiens programmed cell death 5 (PDCD5), mRNA	Homo sapiens golgin-like protein (GLP), mRNA	H,sapiens ART4 gene	60119440F1 NIH MGC 7 Homo saplens cDNA clone IMAGE:3538425 5	Inc28e07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1009284 similar to contains element L1	repeative element;	yassiv. I soares fera liver spiece intres admis sapiers containment.	Mus musculus dynein, axon, heavy chain 11 (Dnahc11), mRNA	Mus musculus dynein, axon, heavy chain 11 (Dnahc11), mRNA	AV723976 HTB Homo sapiens cDNA clone HTBAAG01 5	Homo sapiens golgi autoantigen, golgin subfamily a, 2 (GOLGA2) mRNA	Homo sapíens chromosome 21 segment HS21C027	Homo sapiens chromosome 21 segment HS21C027	H.sapiens DNA for endogenous retroviral like element	Homo sapiens chromosome 21 segment HS21C018	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	Human eosinophil Charcd-Leyden crystal (CLC) protein (lysophospholipase) gene, promoter and exon 1	601467793F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870838 51	RCO-LT0001-150200-032-d11 LT0001 Homo sapiens cDNA	Is56a01.x1 NCI_CGAP_Kid8 Homo sapiens oDNA clone IMAGE:2232552 3'	MR0-HT0923-190800-201-a02 HT0923 Homo sapiens cDNA	as87f12.r1 Strategene fetal retina 937202 Homo sapiens cDNA clone IMAGE:838319 5' similar to	INCUITATION OF A MANUAL	Xp72a03.X1 NCI_CGAP_OV40 Homo sapiens convenients in NASE.X745eee 5	xp72g03.xf NCI_CGAP_Ov40 Homo sapiens cUNA done IMAGE:2/43866 3	Homo sapiens calcium channel, voltage-dependent, alpha 11 subunit (CACNATI), mikiwa Basanganga Nilu MCC 44 Usang samilan sabiba shara NAS 62 sapiensi K	601284360F1 NIM_MGC_44 name septens curva ciane invage, socores o
	Top Hit Database Source	NT	IN	LN	LN	TN	L	NT.	LN	LΝ	EST HUMAN			EST_HUMAN	N	NT	EST_HUMAN	٦	NT	N	된	Ā	F	Ę	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	1000	ESI HUMAN	EST_HUMAN	EST HUMAN	닏	EST HOMAN
	Top Hit Acession No.	5.0E-45 Y18933.1	AB022318.1	5.0E-45 AB022318.1	11496268 NT	11496268 NT	11418704 NT	4759223 NT	8923698 NT	4.0E-45 X95826.1	15 BE 265622.1		15 AA226220.1		6753651 NT	6753651 NT	3.0E-45 AV723976.1	4758451 NT	15 AL163227.2	15 AL 163227.2	3.0E-45 X89211.1	45 AL163218.2	45 AJ243213.1	45 L01665.1	2.0E-45 BE782184.1			45 BE934350.1		45 AA458770.1	AW 270280.1	2.0E-45 AW270280.1	11418157 NT	45 BE389855.1
•	Most Similar (Top) Hit BLAST E Value	5.0E-45	5.0E-45	5.0E-45	5.0E-46	5.0E-45	5.0E-45	5.0E-45	5.0E-45	4.0E-45	4.0E-45		4.0E-45	3.0E-45 T71480.1	3.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	3.0E-45	2.0E-45	2.0E-45	2.0E-45	2.0E-45	2.0E-45	2.0E-45	2.0E-45		2.0E-45	2.0E-45	2.0E-45	20E-45	1.0E-45
	Expression Signal	1.31	0.79	0.79	1.02	1.02	67.0	1.95	2.59	9.58	2.42		0.82	1.35	1.36	1.36	1.4	3.74	13.43	13.43	2.35	2.21	1.22	5.15	1.22	0.78	0.48	18.28		4.16	1.75	1.75	3.93	-1.6
	ORF SEQ ID NO:	31670	31717	31718	31842		34000	34773	37617	26536	27750				31890	31891		34526					28441								37400			
	Exon SEQ ID NO:	18713	18758	18756	18874	18874	20868	21630	24292	13877	15014		21546	16085	18920	18920	21043	21382	22854	22854	25314	15223	15795	19197	20181	21007	22138	25130		23797	24089	24089	ΙI	13185
	Probe SEQ ID NO:	5929	5974	5974	9609	9609	8174	8939	11697	1121	2289		8885	4066	6142	6142	8350	8890	10208	10206	12670	2506	3029	84 82	7510	8314	9485	10705		11129	11488	11488	12711	120

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		T	T	T	T	T	T	T	T	T	T	T	T	Ţ	T	T	T	T	T	T	T	T	Π	T			Γ	T	T	T	Τ	21	7
Single Exon Probes Expressed in Brain	Top Hit Descriptor	601284360F1 NIH MGC 44 Homo saplens cDNA clone IMAGE:3606183 5'	Homo sapiens RAP1A, member of RAS oncodens family (RAP1A), mRNA	Homo sapiens Langerhans cell specific c-type lectin (LANGERIN), mRNA	Human pro-a2 chain of collagen type XI (COL11A2) gene, complete cds	Homo sapiens chromosome 21 open reading frame 1 (C21of4), mRNA	Homo sapiens mRNA for KIAA1591 protein, partial cds	601289116F1 NIH MGC 8 Homo sapiens cDNA clone IMAGE:3619803 5	yr05b02.r1 Soares fetal liver spisen 1NFLS Homo sapiens cDNA clone IMAGE:204383 5	Homo saplens peraxisomal blogenesis factor 14 (PEX14), mRNA	Homo saplens peroxisomal biogenesis factor 14 (PEX14), mRNA	Homo saplens DNA for amyloid precursor protein, complete cds	601511228F1 NIH MGC 71 Homo saplens cDNA clone IMAGE:3912535 F/	Human mRNA for KIAA0299 gene, partial cds	Homo saplens alpha-catenin-like protein (VR22), mRNA	Homo saplens protein kinase C. alpha binding protein (PRKCABP) mRNA	Homo saplens hypothetical protein FLJ20454 (FLJ20454), mRNA	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	Homo sapiens calcium channel, voltage-dependent, alpha 11 subunit (CACNA1), mRNA	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	Homo saplens chromosome 21 segment HS21C009	2822449.5prime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2822449 5'	193708.x1 NCI_CGAP_Gas4 Homo sapiens cDNA cione IMAGE:2132199 3' similar to gb:J00314_ma2_TUBULIN BETA-1. CHAIN (HUMAN);	t3208.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2132199 3' similar to gb;J00314_na2	TOBOLIN BETA-1 CHAIN (HUMAN);	RC5-HT0506-280200-012-C12 HT0506 Homo sapiens cDNA	Rattus norvegicus espin mRNA, complete cds	601277292F1 NIH_MGC_20 Homo sapiens cDNA done IMAGE:3618119 5'	RC4-BT0310-110300-015-f10 BT0310 Homo saplens cDNA	Homo saplens hypothetical protein FLJ10847 (FLJ10847), mRNA	601822835F1 NIH_MGC_77 Homo saplens cDNA clone IMAGE:4042736 5	wm31f08.x1 NCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:24375753' similar to contains MER19.t2 MER19 repetitive element;	
jle Exon Pro	Top Hit Database Source	EST_HUMAN	Ę	Į.	ΙΝ	Į.	N L	EST_HUMAN	EST HUMAN	1	Ę	LN	EST HUMAN	NT	· LV	F	Ę	ト	17	LZ	IN	EST_HUMAN	EST_HUMAN		EST HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	5	EST_HUMAN	EST_HUMAN	
Sing	Top Hit Acession No.	1.0E-45 BE389855.1	4506412	7657290 NT	1.0E-45 U32169.1	8659558 NT	1.0E-45 AB046811.1	-	1.0E-45 H57443.1	11422236 NT	11422236 NT	1.0E-45 D87675.1	45 BE887843.1		19570	11418099 NT	11526291 NT	11418177 NT	11418157 NT	9910293 NT		46 AW246964.1	46 Al433261.1			8.0E-46 BE167244.1		46 BE386165.1 E	46 BE064386.1 E	22708	46 BF105845.1 E	46 AI884381.1 E	
	Most Similar (Top) Hit BLAST E Value	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	1.0E-45	9.0E-46	9.0E-46	9.0E-46	8.0E-46	L	8.0E-46	8.0E-46	7.0E-46	7.0E-46	7.0E-48	7.0E-48	7.0E-46	6.0E-46	
	Expression Signal	2.17	1.38	1.79	7.42	1.38	1.19	5.01	1.04	0.77	22.0	96.0	5.08	1.22	1.33	6.93	11.18	5.28	2.6	1.7	5.86	11.23	13.63	6	13.23	5.69	1.03	3.38	1.33	4.01	1.14	3.90	
	ORF SEQ ID NO:			26569			28962	29804		33752	33753		34861	35272	37651	31125			30991	33958	_	36239	27899	03000	7/200	002200	2/ /00			31692	32170	28207	
	Exon SEQ ID NO:	13185	13245	13906	15866	16240	16315	17178	17411	20825	20625	21197	21709	22100	24327	24592	24708	24711	24969	20821	21224	23024	15162	454.83	20102	7004g	14300	17278	17487	18733	19171	15464	
	Probe SEQ ID NO:	400	460	1151	3101	3483	3560	4442	4677	7930	7930	8505	8019	9422	11734	12087	12263	12269	12680	8127	8532	10378	2443	2773	200	202	7507	4541	4755	5951	6402	2759	

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				_		_	-		_						_	_	_		_	_	_	—,		
Top Hit Descriptor	wm31f08.x1 NCI_CGAP_Ut4 Homo sapiens cDNA clone IMAGE:2437575 3' similar to contains MER19.t2 MER19 repetitive element ;	ta58h10.x1 NCI_CGAP_Kid8 Homo sapiens cDNA clone IMAGE:2232835 3' similar to TR:O60363 O60363 SA GENE.;	xx42e04.X1 NCI_CGAP_UTI Homo saplens cDNA clone IMAGE:27066543' similar to gb:L08069 DNAJ PROTEIN HOMOLOG 2 (HUMAN);	601478409F1 NIH_MGC_68 Hamo sapiens cDNA clone IMAGE:3880895 5'	Homo sapiens chromosome 21 segment HS21C010	7d81g01.x1 Lupski_dorsal_root_ganglion Homo saplens cDNA clone IMAGE:3279408 3'	7d81g01.x1 Lupski_dorsal_root_genglion Homo sapiens cDNA clone IMAGE:3279408 3'	nas38f07.x1 NCI_CGAP_KM11 Homo sapiens cDNA clone IMAGE:3258757 3' similar to TR:O75202 O75202 HOMOLOG OF RAT KIDNEY-SPECIFIC :	602021164F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4156670 5'	QV4-ST0212-120100-075-f09 ST0212 Hamo sepiens cDNA	Z62c08.s1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:726926 3'	no54e09.s1 NCI_CGAP_SS1 Homo eaplens cDNA clone IMAGE:1104520 3' similar to gb:X53741_ma1 FIBULIN-1, ISOFORM A PRECURSOR (HUMAN);	hIB6c03.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3008836 3' similar to gb:X14008_ma1 LYSOZYME C PRECURSOR (HUMAN);contains element MER37 repetitive element ;	hl86c03.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3008836 3' similar to gb:X14008_na1 LYSOZYME C PRECURSOR (HUMAN);contains element MER37 repetitive element;	Human endogenous retrovirus RTVL-H2	Homo sapiens mRNA for KIAA0622 protein, partial cds	Homo sapiens mRNA for KIAA0622 protein, partial cds	Human Ig germline gamma-3 heavy-chain gene V region, partial cds	Human ig germilne gamma-3 heavy-chain gene V region, partial cds	Homo sapiens DNA for Human P2XM, complete cds	Homo sapiens solute carrier family 35 (CMP-stalic acid transporter), member 1 (SLC35A1), mRNA	Homo sapiens VAMP-essociated 33 kDa protein mRNA, complete cds	Homo sapiens mitogen-activated protein kinase kinase kinase kinase 3 (MAP4K3), mRNA	H.sapiens ig kambda light chain variable region gene (7c.11.2) germline; ig-Light-Lambda; VLambda
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	LΝ	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	Z	N	N L	ΡN	NT	۲	ΙN	NT	N	LΝ
Top Hit Acession No.	46 A1884381.1	46 A1635448.1	46 AW513244.1	6.0E-46 BE784971.1	5.0E-46 AL163210.2	BE677194.1	5.0E-46 BE677194.1	46 BF590442.1	5.0E-46 BF347229.1	46 AW 582253.1	46 AA398381.1	46 AA601143.1	46 AW 770544.1	48 AW 770544.1	4.0E-46 M18048.1	46 AB014522.1	4.0E-46 AB014522.1	4.0E-46 M36852.1	46 M36852.1	46 AB002059.1	5453620 NT	46 AF160212.1	4506376 NT	46 273660.1
Most Similar (Top) Hit BLAST E Value	6.0E-46	6.0E-46	6.0E-46	6.0E-46	5.0E-46	5.0E-46	5.0E-46	5.0E-46	5.0E-46	5.0E-46	5.0E-46	4.0E-46	4.0E-46	4.0E-48	4.0E-48	4.0E-46	4.0E-46	4.0E-46	4.0E-46	4.0E-46	3.0E-46	3.0E-48	3.0E-46	3.0E-46
Expression Signal	3.99	10.94	0.72	2.04	8.9	1.07	1.07	1.86	4.20	0.62	0.47	1.4	6.86		2.62	1.04	2.	2.43	2.43	2.71	6.0	0.95	0.79	1.2
ORF SEQ ID NO:	28208	<u> </u>	}			28929	28930			L			27140	27141									29735	
SEQ ID NO:	15484	1		1	13012	16275	•	19398	İ	ı	1	i .	14442	14442	L			1				15150	17100	17456
Probe SEQ ID NO:	2759	8038	7116	11364	198	3519	3519	9839	6842	6995	9515	626	1699	1689	2743	4384	4384	6350	5350	12513	2155	2429	4362	4724

Page 294 of 536 Table 4 Single Exon Probes Fynroscod is

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	H. sepiens ig lambda light chain variable recion gene (7c. 11.2) nerming in Light-Lambda. VI. and de Vienne de Vienn	wj49c04.x1 NCI_CGAP_Lu19 Homo sapiens cDNA clone IMAGE:2406150 3' similar to contains THR.b2 THR repetitive element:	Human mRNA for KIAA0061 gene, partial cds	ne08a08.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:880408 3' similar to contains THR.b2 THR repetitive element:	2/27a11.s1 Soares fetal liver spleen 1NFLS S1 Homo sabiens cDNA close (MACE: 73480a.2)	Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) cenes. complete CRS	Zt89602.r1 Soares_tests_NHT Homo sapiens cDNA clone IMAGE:726650 5' similar to SW:RSP1_MOUSE O01730 RSP-1 PROTEIN	Mus musculus sperm foil assembled match (Stan.)	60144513751 NIH MGC 65 Homo contact Child Act 114 OF COLORS III	v/32d01.r1 Scares fetal liver solden 1NET S. Homo sontone CAIA Alexa 1974 CE Concess of	Xq78h03.X1 NCI CGAP Lu34 Homo sepiens cDNA clone IMAGE-2768789 31	Homo saplens cell division cycle 10 (homolymus to CDC) and sample of division cycle 10 (homolymus to CDC) an	Homo sapiens KIAA0555 cane product / KIAA0555, mDNA	Homo sapiens KIAA0555 gene product (KIAA0555), mRNA	EST390625 MAGE resequences. MAGP Home sealers CDNA	EST48b095 WATM1 Homo seplens cDNA clone 48h095	np78b02.s1 NCi_CGAP_P/2 Homo saplens cDNA clone IMAGE:1132395 similar to gb:X79717 H.sapleno MT-1I mRNA, (HUMAN):	Homo saplens mRNA for KIAA0980 protein partial one	7092b01x/ NCI CGAP OVIB Homo septems cDNA close IMA CE 3844276 21	Homo sapiens centaurin-elohe 2 protein (HSA272405) mBNA	Homo saplens centaurin-alpha 2 protein (HSA272195) mRNA	7092b01.x1 NCI_CGAP_Ov18 Homo saciens cDNA clone IMAGE:3842705.3	df60e03.yf Morton Fetal Cochlea Homo seplens cDNA clone IMA CF-2488844 #	df60e03.y1 Morton Fetal Cochlea Homo sanians cDNA clone IMAGE-248agay #	X02072264F1 NCI CGAP BIR67 Homo saplens cDNA clone IMAGE 4248408 F1	602072264F1 NCI CGAP Bm67 Homo saciens cDNA clone IMAGE 4215398 F	AV715377 DCB Homo sapiens cDNA clone DCBAIE03 5'	Homo sapiens Xq pseudoautosomal region; segment 1/2
gle Exon Pro	Top Hit Datebase Source	۲N	EST HUMAN	LN	EST HUMAN	EST_HUMAN	Į.	EST HUMAN	L	EST HUMAN	EST HUMAN	EST_HUMAN	Z L	L'A	L'Z	EST HUMAN	1.	EST_HUMAN	Į,	EST HUMAN		İ	T HUMAN	EST_HUMAN	T	EST_HUMAN	1	T_HUMAN	IN
Sin	Top Hit Acession No.	46 273660.1	46 AI831462.1	46 D31765.1	46 AA468646.1	46 AA678246.1	46 U78027.1	46 AA399286.1	0569	2.0E-46 BE869151.1	T	4.1	2694	7682177 NT	7662177 NT	6.1	Γ	7:	IG AB023197.1	1.0E-46 BF194707.1	23762	8923762 NT	1.0E-46 BF194707.1	1.0E-46 AW023178.1 E	1.0E-46 AW023178.1	1.0E-46 BF531102.1	1.0E-46 BF531102.1 E		7 AJ271735.1 N
	Most Similar (Top) Hit BLAST E Value	3.0E-46	3.0E-46	3.0E-46	2.0E-46		2.0E-46	2.0E-46	2.0E-46	2.0E-46	2.0E-46 H48391.1	2.0E-46	1.0E-46	1.0E-46	1.0E-48	1.0E-46	1.0E-46 H97330.1	1.0E-46 A	1.0E-46 A	1.0E-46 B	1.0E-46	1.0E-46	1.0E-46 B	1.0E-48 A	1.0E-48 A	1.0E-46 B	1.0E-46 B	1.0E-46 A	9.0E-47 A.
	Expression Signal	1.2	7.59	2.19	7.64	1.55	3.43	1.2	7.67	1.46	1.5	3.38	7.67	1.23	1.23	3.44	3.06	4.55	3.17	88.8	6.14	6.14	5.27	1.53	1.53	2.28	2.28	2.37	6.18
	ORF SEQ ID NO:	30092	34483	37474	26255		27070	30258	33142			30902	26628	26999	27000	27745	27857	28654		31322	31617	31618	31322	37665	37666	31115	31116		1
	Exon SEQ ID NO:	17456		24163		14301	14383	17645	20064	20663	25179	25170	13961	14313	14313	15005	15120	16005	17549	18409	25080	25080	18409	24338	24338	24564	24564	25032	13522
	Probe SEQ ID NO:	4724	8647	11564	817	1554	1637	4917	7384	7968	12257	12575	1211	1566	1566	2279	2399	3243	4818	5613	2888	888	10770	11747	11747	120 44	- 120 44 44	12778	750

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					Signio	201111000	
Probe SEQ ID S NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
4879	17606	30229	3.02	9.0E-4	7 AW770928.1	EST_HUMAN	hi93e04.x1 NCL_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3009534 3' similar to TR:O75703 O75703 HYPOTHETICAL 12.4 KD PROTEIN.;
6284	19057		9.0		11425439 NT	NT.	Homo saplens zinc finger protein ZNF288 (ZNF286), mRNA
12531	25270	30725	2	9.0E-47	11417966 NT	TN	Homo saplens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mRNA
1801	14541	27262				LN.	Homo sapiens HLA-C gene, exon 5, Individual 19323
1801	14541	27253	6.88		8.0E-47 Y18536.1	NT	Homo sapiens HLA-C gene, exon 5, Individual 19323
2722	15429	28167	1.04	8.0E-47	5453955 NT	LΝ	Homo sapiens protein phosphatase 2, regulatory subunit B (B56), epsilon isoform (PPP2R5E) mRNA
3024	15790			8.0E-4		NT	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22, segment 3/3
3613	16366	58009	0.68	8.0E-4		NT	Homo sapiens mRNA for GCK family kinase MINK-2, complete cds
3613	16366	29010				NT	Homo sapiens mRNA for GCK family kinase MINK-2, complete cds
12604	25169		1.38	7.0E-4	7 AV683284.1	EST_HUMAN	AV683284 GKC Homo sapiens cDNA clone GKCASH11 5
2550	15285	28000	1.66	6.0E-4	.2	NT	Homo sapiens chromosome 21 segment HS21C046
8592	21284	34423	0.49	6.0E-4		EST_HUMAN	HSU77054 Human Homo saplens cDNA clone N7
9176	21846		6.76	6.0E4	7 AISS5189.1	EST_HUMAN	t298h02.x1 NCI_CGAP_Kid11 Homo sepiens cDNA done IMAGE:2296659 3'
9612	22265	35450	0.68		6.0E-47 AB042824.1	LN	Homo sapiens RECQL5 beta mRNA for DNA helicase recQ5 beta, complete cds
9612	22265	35451	89.0		6.0E-47 AB042824.1	NT	Homo sapiens RECQL5 beta mRNA for DNA helicase recQ5 beta, complete cds
6482	19249				11423972 NT	INT	Homo saplens CDC37 (cell division cycle 37, S. cerevisiae, homolog) (CDC37), mRNA
10696	23387		6.27		5.0E-47 M78590.1	EST_HUMAN	EST00738 Fetal brain, Stratagene (cat#836206) Homo sapiens cDNA clone HFBCF07
1379	14128	26801	62.8	4.0E-47	4557556 NT		Homo sapiens E1A binding protein p300 (EP300) mRNA
6733	19567	32599	1.9			EST_HUMAN	MR4-TN0108-280800-201-d04 TN0108 Homo sapiens cDNA
8379	21072	34210	2.42			EST_HUMAN	601280486F1 NIH_MGC_39 Homo saplens cDNA clone IMAGE:3622437 6
8379	21072		2.42	4.0E-4	BE616483.1	EST_HUMAN	601280486F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3622437 5
8516	21208	34351	0.61	4.0E-4	7 AW993777.1	EST HUMAN	RC3-BN0034-220300-015-f05 BN0034 Homo sapiens cDNA
1,63,6	2420		283		4 0E-47 AW 515509 1	FST HUMAN	.xx66b07.xd NCI_CGAP_Lym12 Homo sapiens cDNA done IMAGE:2848597 3' similar to SW:IN16_MOUSE Q64252 VIRAL INTEGRATION SITE PROTEIN INT-6. [1];
3 5	42245	25050	205		7 BF907634 1	EST HUMAN	601497639F1 NIH MGC 70 Homo saplens cDNA done IMAGE:3899721 5'
3 2	13315		2 05	3.0E-4	7 BE907634.1	EST HUMAN	601497639F1 NIH_MGC_70 Homo sapiens cDNA done IMAGE:3899721 5'
8	13571			3.0E	17 N57483.1	EST_HUMAN	yy54b04.s1 Soares_multiple_solerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:277327 3'
924	13691	26355		3.0E	47 AL163284.2	٦	Homo sapiens chromosome 21 segment HS21C084
3288	16058	28707	0.79	3.0E	4504116 NT	Ψ	Homo sapiens glutamate receptor, lonotropic, kalnate 1 (GRIK1) mRNA
3948	16698			3.0E~	3.0E-47 U93181.1	Ā	Homo sapiens nuclear dual-specificity phosphatase (SBF1) mRNA, partial cds
4329	17068	29698	1.32	3.0E~	17 M12959.1	뒫	Human T-cell receptor active alpha-chain mRNA from JM cell line, complete cds
5922	18707	31659	5.41	3.0E-	17 AW 408800.1	EST_HUMAN	UI-HF-BM0-adx-d-07-0-UI.11 NIH_MGC_38 Homo sapiens cDNA clone IMAGE:3083205 5

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		T	T		T	T	T	Т	Т	T	T	T	T	T	Т	Т	Т	T	Т	T	T	Т	T	Т	T	7	-	T	Т	T	Т	7
oligie Exoli Flobes Expressed in Brain	Top Hit Descriptor	UI-HF-BM0-sdx-d-07-04.11 r1 NIH MGC 38 Homo septiens churc stone 1886 CE 3083200 E1	Idh04807.xt Scares NET CRC St Homo sentens chala class (MA OE) 442-420	EST375869 MAGE resequences. MAGH Homo sentens CDNA	EST375869 MAGE resequences, MAGH Homo sentions CDNA	Homo saplens myosin phosohatase, fardet suhunit 2 (MYPT2), mBNA	Homo sablens chromosome 21 section HS21Chno	Homo sapiens chromosome 21 segment HS21Chno	WQ96b02.x1 NCI CGAP GC8 Home seniens cONA clone INAGE: 2470884 31	Homo sapiens KIAA0426 gene product (KIAA0428), mRNA	ng43h12.s1 NCI CGAP Co3 Home sentions CDNA / man MACE: 027207.31	Homo saplens ring finger profein (C3HC4 type) 8 (RNFR) mBNA	nf23q07.s1 NCI CGAP Pr1 Homo sepiens cDNA clane IMAGE:01/4852	M23g07.s1 NCI CGAP Pr1 Homo sanions cDNA clans IMA CE:04.4852	Homo sablens Rev/Rex activation domain hinding proteing-related (IDAID) within	EST377239 MAGE resemblences MAGI Home seniors CNA	Homo seplens requisitor of G-powering stoneting & perjent form /DOSE)	801463832F1 NIH MGC 67 Home samens cDNA clave IMAGE 3827407 F1	601463932F1 NIH MGC 67 Home saniens cDNA clans IMAGE 3827497 5	Homo saplens 5-hydroxytrystamine 1D recentor pseudocene with an All repeat incedion	Homo saplens DNA for amyloid precursor protein, complete cda	Homo saplens DNA for amyloid precursor protein, complete cds	Homo saplens SPH-binding factor mRNA, partial cds	Homo saplens BTG family, member 3 (BTG3), mRNA	Human tyrosine kinase receptor (axl) mRNA, complete cds	y92e08.s1 Soares infant brain 1NIB Home sapiens cDNA done IMAGE:29966 3' similar to contains OFR	repetitive element;	qp99h03.x1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:1831189.31	RC3-ST0197-130400-017-h02 ST0197 Homo saplens cDNA	et19e06.x1 Barstead sorta HPLRB6 Homo sepiens cDNA clone IMAGE:2355586 3' similar to gb:M22895 RAS-RELATED PROTEIN RAP-1a /HIMANN	hi84e11.x1 Soeres_NFL_TGBC_STATEMENT CONA clone IMAGE:2978972 3' similar to gb:M26326 KERATIN. TYPE I CYTOSKELETAL 18 (HIMAN)	Papio hamadryas alcohol dehydronenase close I (ADH) come El conica
פום באסוו הוס	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N.	NT	N-	EST HUMAN	N.	EST HUMAN	¥	EST HUMAN	EST HUMAN	Z	EST HUMAN	LN	EST HUMAN	EST HUMAN	NT L	N-I	LN	N-	Ę	NT				EST_HUMAN			NT
	Top Hit Acession No.	-47 AW 408800.1	3.0E-47 AI222413.1	3.0E-47 AW963796.1	3.0E-47 AW963796.1	4505318 NT	2.0E-47 AL163209.2			7662109 NT	AA524514.1	-47 4504886 NT	-47 AA569592.1	-47 AA569592.1	4648	-47 AW965166.1	Γ	Γ		47 L09731.1	47 D87675.1	47 D87675.1	47 AF071771.1	28138	47 M76125.1				47 AW813906.1		85.	
	Most Similar (Top) Hit BLAST E Value	3.0E-47	3.0E-47	3.0E-47	3.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47	2.0E-47			2.0E-47		2.0E-47	1.0E-47	1.0E-47 /	1.0E-47		1.0E-47 L
	Expression Signal	5.41	1.78	0.71	0.71	1.61	2.69	2.69	0.98	1.75	3.41	2	1.5	1.5	1.66	1.3	1.12	1.23	1.23	1.43	1.92	1.92	1.67	0.77	1.27	į	1.75	7.35	1.96	6.79	0.58	2.28
	ORF SEQ ID NO:	31660		34569	34570		26377	26378			27111	29677	29722	29723	28832	30121	31411	31615	31616		33685	33686	34452	35222	36451		30709	26804	30347	32441		36112
	Exon SEQ ID NO:	18707	19236	21424	21424	12958	13713					17052	17090	17090	17206	17493	18490	18673	18673	25118	20559	20669	21310	22051	23218	0,000	20312	14131	17738	19428	21459	22802
	Probe SEQ ID NO:	5922	6469	8732	8732	143	947	947	1560	1588	1673	4313	4351	4351	4471	4761	9699	5887	5887	7598	7864	7864	8618	888	11451	-	17071	202	हुं	6944	8767	10254

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Top Hit Descriptor	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	CM2-MT0100-310700-280-f05 MT0100 Homo sapiens cDNA	601511714F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3913106 5	601511714F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3913106 6	at75h09.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:2377889 3' similar to 1K:060844 060844 HOMOLOG OF RAT ZYMOGEN GRANULE MEMBRANE PROTEIN ;	AU123240 NT2RM1 Homo sapiens cDNA clone NT2RM1000978 5	601310478F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3632083 5	Homo sapiens aminoacylase 1 (ACY1), mRNA	Homo saplens aminoacylase 1 (ACY1), mRNA	hk@1b03.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:3001133 3' sImilar to gb:X64707 BREAST BASIC CONSERVED PROTEIN 1 (HUMAN);	hk81b03.x1 NCI_COAP_Lym12 Homo saplens cDNA clone IMAGE:3001133 3' similar to gb:X84707	March 1 (1997) 1 (199	Homo sapients giutamate receptor, lonouopic, namate i (Oninci) inninci	Iomo sapiens mining lor NAA 1209 protein, partei cus	Homo sapiens mRNA for KIAA 1209 protein, partial cds	Homo sapiens tousled-like kinase 1 (TLK1), mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo sapiens histidyl-tRNA synthetase (HARS), mRNA	Homo sapiens mRNA for AIE-75, complete cds	Homo sapiens BMX non-receptor tyrosine kinase (BMX), mRNA	Homo sapiens mRNA for KIAA1624 protein, partial cds	Homo sapiens mRNA for KIAA1624 protein, partial cds	Homo sapiens putative oncogene protein mRNA, partial cds	Homo sapiens hypothetical protein FLJ11006 (FLJ11006), mRNA	zq45b06.s1 Stratagene hNT neuron (#937233) Homo sapiens cDNA done IMAGE:632827 3' similar to	contains Alu repetitive element,	Homo sapiens phosphodiesterase 1A, calmodulin-dependent (PDE1A) mKNA	RC4-BT0311-141199-011-h06 BT0311 Homo saplens cDNA	MR0-NT0039-010500-002-f08 NT0039 Homo saplens cDNA	tu47a02.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2254154 3	AV690964 GKC Hamo septens aDNA clane GKCDRE125
Top Hit Database Source	TN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	IN	EST HUMAN	INAMI IL FOR	NAMOL DE	Į.	LN-	L L	NT	NT	LΝ	NT	NT	NT	NT	NT	LΝ		EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN
Top Hit Acession No.	AF223391.1	BF359947.1	BE888196.1	BE888196.1	A1833168.1	AU123240.1	BE393813.1	4501900 NT	4501900 NT	AW 768477.1	4 22 4 0 0 2 14 1	AW (004/ /.)	4504116INT	AB033035.1	AB033035.1	6912719 NT	5730038 NT	11416831 NT	AB006955.1	6.0E-48 11420995 NT	B AB046844.1	3 AB046844.1	8 AF026816.1	11427428 NT		6.0E-48 AA189080.1	4826891 NT	5.0E-48 BE064410.1	5.0E-48 AW890299.1	8 AI620420.1	8 AV690964.1
Most Similar (Top) Hit BLAST E Value	9.0E-48		9.0E-48					8.0E-48	8.0E-48		_	-7	8.0E-48	7.0E-48	7.0E-48		7.0E-48	7.0E-48	6.0E-48	6.0E-48	6.0E-4	6.0E-4	6.0E-4	6.0E-48			5.0E-48			4.0E-4	3.0E-48
Expression Signal	3.03	0.82	0.86	0.86	0.66	0.64	3.09	1.44	1.7	4.38			0.79	1.27	17.09	86.0	3.89	27.21				0.58				3.2	1.94				1.27
ORF SEQ ID NO:	27041	09882	31289	l	31754	31877	37002			28541			29302			26916							34866			35425	28713	١.			26786
Exen SEQ ID NO:	14353	L	<u> </u>	ı	1			L						13264	13264					1	١	1	L	L	┖	22241				ı	1 1
Probe SEQ ID NO:	1607	3544	5594	5594	8010	6131	11060	1228	1229	3432		3132	3911	478	479	1483	1634	6460	5967	8889	7365	7365	9022	9441		8896	3304	8474	11603	10878	1364

Page 298 of 536 Table 4 Single Exon Probase Formsond :-

Page 299 of 536 Table 4 Single Exon Probes Expressed in Brain

Single Extri Frodes Expressed in brain	Top Hit Descriptor	td17c01.x1 NCI_CGAP_Co16 Homo saplens cDNA clone IMAGE:2075904 3' similar to TR:014588 014588 SIMILARITY TO U73941;	Homo sapiens NF2 gene	Homo sapiens mRNA for KIAA1071 protein, partial cds	Homo sapiens mRNA for KIAA1071 protein, partial cds	Homo sapiens huntingtin (Huntington disease) (HD) mRNA	Homo saplens mitogen-activated protein kinase kinase kinase 13 (MAP3K13), mRNA	Homo sapiens mitogen-activated protein kinase kinase kinase 13 (MAP3K13), mRNA	Homo sapiens Chediak-Higashl syndrome 1 (CHS1) mRNA	Homo sapiens mRNA for KIAA1245 protein, partial cds	601888086F1 NIH_MGC_17 Hamo sapiens cDNA clone IMAGE:4122119 5'	Homo saplens B cell linker protein (SLP65), mRNA	Homo saplens B cell linker protein (SLP65), mRNA	Mus musculus MysPDZ mRNA for myosin containing PDZ domain, complete cds	Mus musculus T-box 20 (Tbx20), mRNA	Mus musculus T-box 20 (Tbx20), mRNA	Human inositol 1,4,5 trisphosphate receptor type 1 mRNA, partial cds	Homo sapiens gene for activin receptor type IIB, complete cds	Homo saplens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo saplens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens proteasome (prosome, macropain) 28S subunit, ATPase, 4 (PSMC4) mRNA	Homo sapiens chromosome 21 segment HS21 C084	w/25h04.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2356663 3' similar to TR:054923 O54023 RSEC46	DKF2n7620033 s.1 762 (synonym: hmel2) Homo sablens cDNA clone DKFZn7620033 3'	w/25h04 v1 Soares NFL T GBC S1 Homo seniens cDNA clone IMAGE:2356663 3' similar to TR:054923	064823 RSEC15.;	ba55g05.x1 NIH_MGC_10 Homo sepiens cDNA clone IMAGE:2900504.3' similar to gb:X17206.40S DIBOSOMAI PROTEIN SA (HIIMAN): Ar-MADK33 Marise II Rev3 notein mRNA from a renetitive element	ocmplete (MOUSE);	DKFZp761A138_s1 761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761A138 3
ום בייחוו ביחחם	Top Hit Database Source	EST_HUMAN SI	NT TN)H	NT TN					NT TN	EST_HUMAN 60			ĮŃ LN			Ή LN	IN IN							NT H	N C	Т	Т	EST_HUMAN O		EST_HUMAN \alpha	EST_HUMAN D
BILLO	Top Hit Acession No.	48 AI889077.1	48 Y18000.1		48 AB028994.1	4755137 NT	4758695 NT	4758695 NT	4502838 NT			11429808 NT	11429808 NT	-49 AB026497.1	10048417 NT	10048417 NT	49 U23850.1	Ξ	5729990 NT	5729990 NT	5729990 NT	5729990 NT	5728990 NT	5729990 NT	49 AL163284.2		7.0E 40 At 420037 4	Τ	-49 A3807191.1		6.0E-49 AW731740.1	6.0E-49 AL162091.1
	Most Similar (Top) Hit BLAST E Value	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.05-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	8.0E-49	8.0E-49	8.0E-49	8.0E-49	8.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49	7.0E-49	07 30 2	7.00-10	01-101-1	7.0E-49		6.0E-49	6.0E-49
	Expression Signal	1.17	46.0	0.71	0.71	2.52	92.0	0.76	0.84	6.4	5.33	4.08	4.08	1.13	3.43	3.43	3.17	1.15	-	-	1.73	1.73	2.94	2.94	3.4	77.0	- 4	-	0.67		12.12	1.27
	ORF SEQ ID NO:	31947		32266	32267	32913	34566	34567	34966	35004	35314	36127	36128	27461	31704	31705	34026	35733	25814	25815	25814	25815	25814	25815	26615	0000	30002	Cions	30862		25646	29464
	Exan SEQ ID NO:	18971	19178	19265	19265	19844	21422	21422	21801	21838	22134	22917	22917	14737	18744	18744	20888	22539	13171	13171	13171	13171	13171	13171	13951	5,6,	101/3	300	18173		13005	16837
	Probe SEQ ID NO:	6195	6407	9200	9200	7157	8730	8730	9113	9168	9481	10269	10269	2002	2962	2962	8194	6886	135	135	ğ	38	385	385	1199	G. C.	3 8	3	5718	<u> </u>	182	4085

Page 300 of 536 Table 4 Single Exon Probes Expressed in Braln

		Т	0	Т	Т	Т	Ţ	Т	Т	T	Т	I	T	T	T	Т	T	7	Т	Т	T	T	_	_	Τ_	_	•
The court is provided in the court is the court in the co	Top Hit Descriptor	AU140742 PLACE4 Homo saplens cDNA clone PLACE4000148 5'	hd44e02.x1 Soares, NFL_T_GBC_S1 Hamo sapiens cDNA clone IMAGE:2912378 3' similar to TR:095636 O95636 CAMP-REGULATED GUANINE NUCLEOTIDE EXCHANGE FACTOR II	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	UI-H-Bi3-alo-a-05-0-UI s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:3068048 3'	EST77525 Pancreas tumor III Homo sapiens cDNA 6' end	EST77525 Pancreas tumor III Homo sapiens cDNA 6 end	2/29c08.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo septens cDNA clone IMAGE:4616943	Homo sapiens chromosome 21 segment HS21C010	Homo saplens chromosome 21 segment HS21C010	2p29c07.r1 Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone IMAGE:610860 5' similar to TR:0233226 G233226 RTVL-H PROTEIN :contains LTR7 t3 LTR7 repetitive element	Homo sapiens putative tumor suppressor ST13 (ST13) mRNA, complete ods	Homo sapiens similar to ribosomal protein S27 (metallopanstimulin 1) (H. sapiens) (LOC63362), mRNA	XI08b01.x1 NCI_CGAP_Ut4 Homo saplens cDNA clone IMAGE:2075583 3' similar to WP:B0350.2B CE06703 :	Homo sapiens UDP-N-acety-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 8 (GalNAc-T8) (GALNT8), mRNA	Homo explens UDP-N-acetyl-alpha-D-galactosemine:polypeptide N-acetylgalactoseminyltransferace 8 (GaINAc-T8) (GALNT8), mRNA	Homo sapiens KIAA0623 gene product (KIAA0623), mRNA	Homo sapiens copine III (CPNE3), mRNA	Homo sapiens copine III (CPNE3), mRNA	Z90f05.r1 NCI_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:682977 67	Homo sapiens glutathlone S-transferase theta 2 (GSTT2) and glutathlone S-transferase theta 1 (GSTT1)	genes, complete cds	H.sapiens mRNA for acetyl-CoA carboxylase.	ze31c05.r1 Soares retina N2b4HR Homo septens cDNA clone IMAGE:360884 5' similar to contains L1.13 L1 repetitive element;	Human type IV colladen (COL4AB) dene, exon 40	
ממו ווומשי מומי	Top Hit Database Source	EST_HUMAN	EST HUMAN			T HUMAN	EST_HUMAN	Ι'''	Г	Г			Į.		EST HUMAN						EST_HUMAN 2			Į.	EST HUMAN IN	T	
5	Top Hit Acession No.	19 AU140742.1	19 AW611225.1	9910293 NT	9910293 NT	19 AW452218.1	6.0E-49 AA368556.1	6.0E-49 AA366556.1	6.0E-49 AA707567.1	AL163210.2	5.0E-49 AL163210.2		5.0E-49 U17714.1	6355	4.0E-49 AW189533.1	11525737 NT	11626737 NT	7662209 NT	11425374 NT	11425374 NT	9 AA210798.1		4.0E-49 AF240786.1		3.0E-49 AA016131.1		
	Most Similar (Top) Hit BLAST E Value	6.0E-49	6.0E-49	6.0E-49	6.0E-49	6.0E-49	6.0E-49	6.0E-49	6.0E-49	5.0E-49	5.0E-49	5.0E-49	5.0E-49	6.0E-49	4.0E-49	4.0E-49	4.0E-49	4.0E-49	4.0E-49	4.0E-49	4.0E-49		4.0E-49/	3.0E-4912	3.0E-49/	3.0E-49 U46999 1	
	Expression Signal	0.94	69:0	0.45	0.45	2.6	2.6	2.6	2.03	2	7	3.49	4.25	2.64	47.84	0.95	0.95	6.0	0.47	0.47	4.21		4.1	1.73	9.	2.33	
	ORF SEQ ID NO:	32109	33076	35478		37202	37579	37580		26117	26118	27235	28201	28678	25927	32930	32931	33488	34604	34605				25961		30264	
	Exon SEQ ID NO:	19119	19997			23910	24257	24257	25151		13470	14527	15459	16029	13298	19858	19858	20374	21466	21455	25368	- 6	24730	13330	15364	17851	
	Probe SEQ ID NO:	6349	7314	9633	9633	11248	11681	11661	12362	695	969	1786	2764	3267	512	7172	7172	7710	8763	8763	12221		12306	\$	2654	4923	

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Probe SEQ ID NO: 11272 11272 11272 12316 1546 1794 1794 1546 1717 1717 1717 1717 1717 1717 1717 1717 1717 1712 17200 172	Exon SEQ ID NO: 17426 17426 17426 17426 17426 17426 17426 17534 18080 19805 19805 19805 19805 19805 19805	ORF SEQ ID NO: 30110 30110 30118 30110 30118 30737 31734 31734 31734 32860 32860	Signal 1.98 2.94 1.06 0.68 0.68 0.74 4.82 2.07 2.07 2.07 2.07 2.07 2.07 2.07 2.0	Most Simila (Top) Hit BLASTE Value 3.0E-4 2.0E-4 2.0E-4 1.0E-4 TE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE No. 4STE NO. 4ST	Top Hit Database Source Source Source EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN	Top Hit Descriptor EST42572 Endometrial tumor Homo saplens cDNA 6' end MR3.HT0487-150200-113-g01 HT0487 Home saplens cDNA clone IMAGE:262571 5' OZ88d02.x1 Scares_senescent_fibroblasts_NIN45F Homo saplens cDNA clone IMAGE:262571 5' OZ88d02.x1 Scares_senescent_fibroblasts_NIN45F Homo saplens cDNA clone IMAGE:1682403 3' similar to gb:M31470 RAS-LIKE PROTEIN TC:10 (HUMAN);contains Alu repetitive element; IUH-Bit-ense-d-02-0-UI.s1 NCI_CGAP_Sub8 Homo saplens cDNA clone IMAGE:3088538 3' AV717338 DCB Homo saplens cDNA done DCBALB01 6' EST02558 Fetal brain, Strategene (catta305206) Homo saplens cDNA clone HFBCY50 Homo saplens keratiun 8 (KRT19) mRNA For 145769F1 MIH_MGC_6 Homo saplens cDNA clone IMAGE:3862086 6' For 15769F1 MIH_MGC_6 Homo saplens cDNA clone IMAGE:386208 6' For 15769F1 MIH_MGC_8 Homo saplens cDNA clone IMAGE:386208 6' For 16769F1 MIH_MGC_8 Homo saplens cDNA clone IMAGE:3862088 6' For 18770300 ADB Homo saplens cDNA clone Bit IMAGE:3862088 6' For 18770300 ADB Homo saplens cDNA clone ADBCVD11 6' For 18770300 ADB Homo saplens cDNA clone ADBCVD11 6' For 18770300 ADB Homo saplens cDNA clone IMAGE:3520863 6' For 18770300 ADB Homo saplens cDNA clone IMAGE:3520863 6' For 18770300 ADB Homo saplens cDNA clone IMAGE:3520863 6' For 18770300 ADB Homo saplens cDNA clone IMAGE:3520863 6' For 18770300 ADB Homo saplens cDNA clone IMAGE:3520863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3620863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187703100 ADB Homo saplens cDNA clone IMAGE:3630863 6' For 187	
7977	1 1			1.0E.4	11321580 NT	본	Homo sapiens succinate CoA ligase, GDP-forming, alpha subunit (SUCLG1), mRNA
7977	1				11321580 NT	NT	Homo septens succinate-CoA ligase, GDP-forming, alpha subunit (SUCLG1), mRNA
8575	1 1	П		1.0E-4	9994184 NT	NT	Homo saplens RNA binding motif protein 7 (LOC51120), mRNA
88 8	i	34721	1.29	1.0E.4	9 BE409340.1	EST HUMAN	601300992F1 NIH_MGC_21 Homo sepiens cDNA clone IMAGE:3635398 5' [DKF7-A34D73423 -1 434 (summum: hime3) Homo seniens cDNA clone DKF7-A34D7423 5'
10028	- 1	1			9 AL043129.2 9 AV751477.1	EST HUMAN	DKF2p434D2423_T1 434 (Synonym: mess) fromo sapiens cunk clone DKF2p434D2423 5 AV751477 NPD Hamo sapiens cDNA clone NPDAWE04 5
11281	_L_			1.0E-49	11427366 NT	. 1	Homo saplens brefeldin A-Inhibited guanine nucleotide-exchange protein 1 (BIG1), mRNA
12215	1	j			11418322 NT	NT	Homo sepiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA
4937	ll		1.4		9.0E-50 AF101475.1	뒫	Homo sapiens glycine N-methyltransferase (GNMT) gene, complete cds

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| Top Hit Descriptor | 601176250F1 NIH MGC 17 Home septens CDNA clane MACE-3534588 F | Homo saplens chromosome 21 segment HS21C002 | Homo saplens mRNA for VIP receptor 2 | Homo sapiens mRNA for VIP receptor 2 | Homo sapiens actinin, alpha 1 (ACTN1) mRNA | Homo saplens capping protein (actin filament) muscle Ziline heta (CAPZR) mBNA | Homo saplens hepatocyte growth factor(HGF) gene, expn 18 | Inp62d06.s1 NCI_CGAP_Br2 Home sapiens CDN4 clone IMAGE:1130891 3' similar to gb:J06459 GLUTATHIONE S-TRANSFERASE TESTIS/RDAIN (UI IMAN)
 | QV0-BT0703-280400-211-608 BT0703 Homo sepilens cDNA | IRC6-TN0073-150900-011-A12 TN0073 Home saniens cDNA | RC8-TN0073-150900-011-A12 TN0073 Homo saniens cDNA | ng59e12.s1 NCI_CGAP_Co9 Homo sepiens cDNA clone IMAGE:1148208 3' similar to gb:X68391 60S
RIBOSOMAL PROTEIN L6 (HUMAN): | Wm55g11.x1 NCI CGAP Ut2 Home septems cDNA clane IMAGE 2430008 31 | 601569565F1 NIH MGC 7 Homo sapiens cDNA clone IMAGE:3943677 5
 | ho39h04.x1 NCI_CGAP_Ut1 Homo saplens cDNA clone IMAGE:3039511 3' similar to contains MER29.b3 MER29 repetitive element | EST182775 Jurkat T-cells VI Homo saniens cINA 5' and | EST182775 Jurkat T-cells VI Homo saplens cDNA 5' and | CM0-BT0792-300500-398-b05 BT0792 Homo septens cDNA

 | CM0-BT0792-300500-398-b05 BT0792 Homo sapiens cDNA | n/45h10.s1 NCI_CGAP_Pr4 Homo saplens cDNA clone IMAGE:1043683 similar to contains PTR6.t3 PTR6 repetitive element;
 | no54e09.s1 NCI_CGAP_SS1 Home sapiens cDNA clone IMAGE:1104520 3' similar to gb:X53741_ma1
FIBILIIN-1 ISOEORM & DEFCLIDE OF ALLIMANN. | Homo sablens chromosome 21 serment Li2247049 | Homo sapiens cystein/J-RNA syntheties (CADS) DNA
 | OV1-BT0681-280300-197412 RT0681 Home senions ADNA | Human endodemoria retroduris RTVI -H2 | ob03/06.s1 NCL CGAP Kid3 Homo sepiens cDNA clone IMAGE-1322827.3 | GMYA5 Human cardiac muscle expression library Homo sapiens cDNA cione 4151935 similar to CMYA5 Cardiomyopathy associated gene 5 |
| Top Hit
Database
Source | EST HUMAN | NT TN | LN | NT | NT | K | N. | EST HUMAN
 | EST HUMAN | EST HUMAN | EST HUMAN | EST HUMAN | EST HUMAN | EST_HUMAN
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| Top Hit Acession
No. | BE295758.1 | AL163202.2 | X95097.2 | X95097.2 | 4501890 | | D90334.1 |
 | ŀ | | BF091922.1 | AA627822.1 | |
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 | | T | 10683
 | | | | 3.0E-50 AW755254.1 |
| Most Similar
(Top) Hit
BLAST E
Value | 9.0E-50 | 8.0E-50 | 8.0E-50 | 8.0E-50 | 8.0E-50 | 8.0E-50 | 8.0E-50 | 8.0E-50
 | 7.0E-50 | 7.0E-50 | 7.0E-50 | 7.0E-50 | 7.0E-50 | 8.0E-50
 | | 6.0E-50 | 6.0E-50 | 5.0E-50

 | 5.0E-50 | 5.05-50
 | 4.0E-50 | 4.0E-501/ | 4.0E-50
 | 4.0E-50 | 3.0E-50 | 3.0E-50/ | 3.0E-50 A |
| Expression
Signal | 0.95 | 4.05 | 2.54 | 2.54 | 2.82 | 1.48 | 15.7 | 1.29
 | 0.78 | 1.06 | 1.08 | 0.6 | 7.65 | 0.68
 | 5.67 | 12.6 | 12.6 | 1.1

 | 1,1 | 5.26
 | 1.7 | 0.99 | 0.98
 | 1.95 | 4.13 | 1.24 | 1.14 |
| ORF SEQ
ID NO: | | | 26125 | 26128 | | | · | 37293
 | 26015 | 32643 | 32644 | 32966 | 36584 |
 | | 36846 | 36847 | 27233

 | 27234 |
 | | 28847 | 32018
 | 32888 | - | 28703 | 29124 |
| | | | | | | ı | 14691 | _
 | ı | | 19604 | 19890 | 23347 | 17048
 | 20806 | 23406 | 23406 | 14526

 | 14526 | 21680
 | 13665 | 16197 | 19041
 | 19822 | 14667 | 16054 | 16487 |
| Probe
SEQ (D
NO: | 6310 | 166 | 702 | 702 | 1768 | 2703 | 2833 | 11385
 | 802 | 6687 | 6687 | 7205 | 10856 | 4309
 | 8112 | 10717 | 10717 | 1785

 | 1785 | 8990
 | 897 | 3441 | 8529
 | 7135 | 1831 | 3293 | 3734 |
| | Exon ORF SEQ Expression (Top) Hit Top Hit Acession No. Signal BLASTE No. Source | Exon ORF SEQ Expression Signal (Top) Hit Acession (Top) Hit Acession Signal Top Hit Acession (Top) Hit Acession Signal One Signal One Signal Top Hit Acession Source Source Source Source Source Source Source Adales | Exon ORF SEQ Expression Signal (Top) Hit Acession No: Top Hit Acession Signal Top Hit Top Hit Acession No: Signal Top Hit Top Hit Acession No: Source Source Top Hit Acession Source Source Database Source Source Source Source Source Acession Source Source Acession Source Sour | Exon
SEQ ID
NO: ORF SEQ
ID NO: Expression
Signal (Top) Hit
ID NO: Top Hit
No. Top Hit
Source 26421 0.96 9.0E-50 BE295768.1 EST_HUMAN 12380 25619 4.06 8.0E-50 AL163202.2 NT 13477 26128 2.54 8.0E-50 X95097.2 NT | Exon ORF SEQ Expression Signal (Top) Hit Acession Signal (Top) Hit Acession Signal No. Top Hit Acession Source Native Source | Exon ORF SEQ Expression Signal (Top) Hit Acession Signal (Top) Hit Acession Signal No. Top Hit Acession Source Nation Signal No. Top Hit Acession Source Source Nation Source Source Source Source Source Source Source Nation Source Source Source Source Nation Source Sour | Exon
SEQ ID
ID NO: CAPE SEQ
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ID NO: Expression
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Signal
ID NO: Most Similar
FLASTE
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13477 Top Hit Acession
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A.05 Top Hit Acession
No. Top Hit Acession
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A.163202.2 Top Hit Acession
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A.163203.2 Top Hit Acession
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ID Exon
SEQ ID
NO: ORF SEQ
Signal Expression
Signal (Top) Hit
BLASTE
Value Top Hit
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Source
Source 25421
12980 25619
25618 4.05
4.05
4.05
4.05
8.0E-50 9.0E-50
8.0E-50 BE295789.1
AL163202.2
AL163202.2
NT EST HUMAN 13477 26125
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Table 4
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
2999	19584	32618	1.52	3.0E-50	11421514 NT	TN	Homo sepiens similar to sema domain, immunoglobulin domain (ig), short basic domain, secreted, (semaphorin) 3A (H. sepiens) (LOG63232), mRNA
7644	20214	33314	4.85		AF233436.2	- TN	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1a mRNA, complete cds
7544	20214	33315	4.85			Į.	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP1a mRNA, complete ods
8481	ı	L			6601589 NT	LZ	Homo sapiens ankyrin-like with transmembrane domains 1 (ANKTM1), mRNA
9718	22369		1.21		3.0E-50 AB046818.1	NT	Homo sapiens mRNA for KIAA1598 protein, partial cds
9727			66.0		11418514 NT	NT .	Homo sapiens t-complex 10 (a murine tcp homolog) (TCP10), mRNA
10077			0.47			NT	Mus musculus mRNA for neurobeachin
10415					3.0E-50 AB002297.1	NT	Human mRNA for KIAA0299 gene, partial cds
11045	23715	36984	1.61		36955	LN	Homo sapiens Grb2-associated binder 2 (KIAA0571), mRNA
11441				i	3.0E-50 AJ245621.1	LN	Homo saplens CTL2 gene
760	13532				2.0E-50 AF055066.1	NT	Homo sapiens MHC class 1 region
1057		,	5.67		4657752 NT	ΝŢ	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA
1424			2.25		2.0E-50 AF138303.1	NT	Homo sapiens decorin D mRNA, complete cds, alternatively spliced
69/9	19513		0.59			EST_HUMAN	AU124085 NT2RM2 Homo sapiens cDNA clone NT2RM2001609 5
8215		34044	1.02		2.0E-50 AB038162.1	NT	Homo sapiens TFF gene cluster for trefoil factor, complete cds
8215				. 1		NT	Homo sapiens TFF gene cluster for trefoil factor, complete cds
8355	21048	ŀ				TN	Human HALPHA44 gene for alpha-tubulin, exons 1-3
8356		l				TN	Human HALPHA44 gene for alpha-tubulin, exons 1-3
9784	l I					NT	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA
9784	22435	35642	1.51	2.0E-50	TN 6820188	NT	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA
11660	24256		1.8			TN	Macaca mulatta cyclophilln A mRNA, complete cds
449	13236	25874	1.92		1.0E-50 AL163209.2	NT	Homo saplens chromosome 21 segment HS21C009
2365	15087		87.6			LN	Homo sapiens Xq pseudoautosomal region; segment 1/2
10093	22741	35956	1.57			M	Homo sapiens RGH2 gene, retrovirus-like element
				100	, 2007, 21, 1	1100	hd44e02x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2912378 3' similar to TR:095636
5893				9.0E-5		EST HOMAN	OBBOSS CAMP-KECOLA IEU GUANINE NUCLEO I IDE EXCHANGE PACI UK II.
6130	18908	31876	0.71	9.0E-51	9.0E-51 AA744837.1	EST HUMAN	ny67h03.s1 NCL_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1283381 3'
0577	77.067	24400	38 0		0.05 64 81204464.4	TOT LIMAN	ab23g04.x5 Stratagene lung (#337210) Homo sapiens cDNA clone IMAGE:841686 3' similar to SW:DeM HI IMAN Ondero PROSTATE.SPECIEIC MENARANIFANTICEN :
7/00	- 1				T		THE COLUMN CONTROL OF THE COLUMN COLUMN CONTROL OF THE COLUMN CONTROL OF THE COLUMN CONTROL OF THE COLUMN COLUMN COLUMN CONTROL OF THE COLUMN COLUMN COLUMN COLUMN CONTROL OF THE COLUMN COLUMN COLUMN
9224	21903	35075	1.23		9.0E-51 AA043738.1	ES HOMAN	ZX31QU9.11 Society pregnant uterfus North Homo Repiens CUNA Glotte IMACE:466352 6

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					ושום ביטוו רוט	Single Exort Probes Expressed in Brain
SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
22062	35231	99.0	9.0E-	51 A1791154.1	EST_HUMAN	ab23g04.x5 Stratagene lung (#937210) Homo saplens cDNA clone IMAGE:841686 3' similar to SW:PSM_HUMAN Q04609 PROSTATE:SPECIFIC MEMBRANE ANTIGEN
22062	35232	99.0	9.0E-51	51 AI791154.1	EST HUMAN	ab23g04.x6 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:841686 3' similar to SW:PSM HUMAN Q04609 PROSTATE-SPECIFIC MEMBRANE ANTICEN.
23222			9.0E-51	H89078.1	EST HUMAN	W24g06.r1 Mortan Fetal Cochlea Hamo sanieras cDNA clone IMA GE-25324.0 gr
23222			9.0E-51		EST HUMAN	Iw/24g06.r1 Morton Fetal Cochlea Homo sepiens cDNA clane IMAGE-283210.5
18908		1.43	9.0E-51	AA744837.1	EST HUMAN	IN967h03.s1 NCI CGAP GCB1 Homo sapiens cDNA clone IMAGE 12833210 5
17142	29770			4503932	Ę	Homo saplens glycine amidinoransferase (L-aminine/nycine amidine/mensferase (L-aminine/nycine amidine/nycine)
17142		1,45	8.0E-51	4503932 NT	K	Homo saplens glycine amidinofransferase (L-arginine: glycine amidinofransferase) (OA TA) DNA
17265				8.0E-51 AA610842.1	EST HUMAN	np98e09.s1 NCI_CGAP_Lu1 Homo sapiens cDNA clone IMAGE:1142440 3' similar to gb:X12671_ma1 HETEROGENEOUS NUCLEAR RIBONI ICI ECOPOTEIN 44 441 11.401.
20222	33325		8.0E-51	282	Į.	Homo saplens PDZ-73 ministry (PDZ-73ANY-CC-20)
21939		1.13	8.0E-51		EST HUMAN	AU138590 PLACE1 Homo saniens CDNA class DI ADE1000807 ET
20222	33325	2.02	8.0E-51	8.0E-51 11439687 NT	L	Homo saplens PDZ-73 protein (PDZ-73/NY-CO-38) mRNA
15781	28430	0.9	7.0E-51		EST HUMAN	xn34a03.x1 NCI_CQAP_Kid11 Homo sapiens cDNA clone IMAGE:2696684 3' similar to TR:Q9Z340 Q9Z340 ATYPICAL PKC SPECIFIC RINDING PEDETEIN
16037	28687	1.45	7.0E-51	Γ	EST HUMAN	0V4-NT0028-200400-180-406 NT0028 Home series ability
16888	29519	1.37	7.0E-51	Τ	EST HUMAN	DKFZ043482229 r1 434 (synonym: hips3) Home seniens cDNA clara DKFZ-43482229 F1
16888	28520	1.37	7.0E-51	Τ	Т	DKF20434B2229 11 434 (smonwar: bless) Home septems CDNA close DKF2-40450635 21
17057	29681	2.71	7.0E-51	7.0E-51 AW295603.1	Т	UI-H-BW0-aip-b-05-0-UI-s1 NOI CGAP Subs Homo saniens CDNA chara IMAOE 2720047 21
24281	37603	1.34	7.0E-51			Homo sapiens HSPC331 mRNA, partial cds
14708	27426	4.96	6.0E-51	7657266 NT		Homo sapiens KIAA0929 protein Msx2 intaracting nuclear target (MINT) fromolog (KIAA0929), mRNA
16222	28876	14.73	6.0E-51	7657286INT		Homo saplens KIAA0929 nmlein Mev2 Intercenting minimum services to the minimum services of the minimum services of the minimum services of the
18686	31634	1.56		X01788.1		Human haptoplobin related (Hnr) name even 3
18696	31648	9.85		AF070083.1		Omo sablens mitroren-artiusted protein kinasa kinasa 4 (ARVA)
18696	31649	9.95		Γ		Homo sapiens mitpoen-activated protein kinase kinase 1 (MKKA) anno 2000 4
19580	32615	1.02	6.0E-51	4506736 NT		Homo sapiens ribosomal protein Se kinasa 70km Polyvanitha 4 / PBSeVB4 - BVIA
19536	32564	0.97	6.0E-51	11416751 NT		Homo saplens non-kinasa Cdo42 effector protein SPEC2 (Octagoo) DNA
17945	30540	. 2.2	6.0E-51	11429665 NT		Homo saplens carebral cell adhasion molecula (I OCB1448) mBNA
21725	34878	0.68	6.0E-51	11428525 NT		Homo saplens hypothetical protein FL 111042 (FL 111042), mRNA
21725	34879	0.68	6.0E-51	11428525 NT		Homo saplens hypothetical protein FLJ11042 (FLJ11042) mRNA
J	35419	2.18	6.0E-51	7661535 NT		Homo saplens B9 protein (B9), mRNA

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Single Exon Probes Expressed in Brain

	Top Hit Descriptor	(1) gene, expn 2	aukin 17 receptor (IL17R), mRNA	Homo sapiens protein phosphatase 2, regulatory subunit B (856), albha isoform (PPP2R5A) mRNA	Homo sapiens protein phosphalase 2. regulatory subunit B (R58), alpha is notwern (PDD2DFA) with a	10some 21 segment HS21C003	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	rapping to chamosome X	Homo sapiens 26S proteasome-associated pad1 homolog (POH1) mRNA	A for nucleaparin 155	subunit mRNA, complete cds	subunit mRNA, complete cds	binding motif protein 3 (RBM3), mRNA	tr81c09.xt NCI_CGAP_Part Homo sapiens DNA done IMAGE:2224720 3' similar to gb:M26326 KERATIN, TYPE I CYTOSKELETAL 18 (HUMAN):	lapping to chomosome 22	9847c08.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:53233 5' similar to gb:M14123_cds4 RFTROVIRI IS.REI ATED BOIL DOI VODOTEIN JULIMANIA.	Total mRNA	ancreatic Islets Homo sapiens cDNA 5'	Homo saplens ubiquitin protein ligase E3A (human papilloma virus E8-associated protein, Angelman	mRNA	z/30a05.r1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone IMAGE:664880 5' similar to TR:6233226 G233226 RTVL-H PROTEIN: contains LTR7,13 LTR7 repetitive element:	t27g03.x1 NCI CGAP Kid11 Homo saplens cDNA done IMAGE:2131732 3'	UI-H-B11-adj-d-02-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2716851 3	obs409.x5 NCI_CGAP_Xid5 Homo sapiens cDNA clone IMAGE:1325609 3's similar to SW:NME1_MOUSE P35436 GLITAMATE INMIDALRECEPTOR SUBLINIT EDSUION 1 PDE CUIDSOD.	ob34f09.x5 NCI_CGAP_Kid5 Homo sapiens cDNA clone IMAGE:1325609.3° similar to SW:NME1_MOUSE P34436.01 i TAMATE INAMA INCEDED SUBJECT SUBJECT CONTROL OF THE INAMATE INAMA INCEDED SUBJECT CONTROL OF THE INAMA INCEDIT CONTROL OF THE INAM	601470446F1 NIH MGC 67 Home septens CDNA clone IMAGE-3872563 51	ilycerol kinase lota (DGKI) gene, exon 23
מספת ווו בותווו	Top Hit	Human ankyrin (ANK1) gene, exon 2	Homo saplens interleukin 17 receptor (IL17R), mRNA	lens protein phosphatase 2, regulatory subu	iens protein phosphatase 2. regulatory subu	Homo saplens chromosome 21 segment HS21C003	iens T-cell lymphoma invasion and metastas	Novel human gene mapping to chomosome X	iens 26S proteasome-associated pad1 homo	Homo sapiens mRNA for nucleoporin 155	Human Ku (p70/p80) subunit mRNA, complete cds	Human Ku (p70/p80) subunit mRNA, complete cds	Homo sapiens RNA binding motif protein 3 (RBM3), mRNA	#81c09.x1 NCI_CGAP_Pan1 Homo saplens cDNA cl	Novel human gene mapping to chomosome 22	Soares infant brain 1NIB Homo saplens of	Human hnRNP C2 protein mRNA	la04d06.v1 Human Pancreatic (slets Homo sapiens cDNA 5	ens ubiquitin protein ligase E3A (human par	syndrome) (UBE3A) mRNA	Stratagene NT2 neuronal precursor 83723(3226 G233226 RTVL-H PROTEIN.; contain	NCI CGAP Kid11 Homo saplens cDNA dic	dj-d-02-0-Ul.s1 NCI CGAP Sub3 Homo se	SNCI_CGAP_Kid5 Homo sapiens cDNA clo	NO! CGAP Kid5 Homo sapiens cDNA clo	FT NIM MGC 67 Homo septens CDNA clos	Homo saplens diacyglycerol kinase lota (DGKI) gene, exon 23
2000	Hit	Human an	Homo sap	Homo sap	Homo sap	Homo sap	Homo sapi	Novel hum	Homo sapi	Homo sapi	Human Ku	Human Ku	Homo sapi		Г		Т	Г	Homo sapi	syndrome)		Γ	Г			Т	П
	Top Hit Database Source	μ	PINT	TN 6	FN 6	Ϋ́	TNC	¥	₩.C	Έ	Ä	Ę	3NT	EST HUMAN	Į.	H HAAN	N-	EST HUMAN			EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	ENT EN IMAGN	EST HUMAN	Ę
;	Top Hit Acession No.	6.0E-51 U50093.1	11528289 NT	5453949 NT	5453949 NT	5.0E-51 AL163203.2	4507500 NT	5.0E-51 AL133204.1	5031980 NT	5.0E-51 AJ007558.1	M30938.1	5.0E-51 M30938.1	5803136 NT	AI587348.1	3.0E-51 AL 159142.1	3.0F-51 R15914 +	M29063.1	3.0E-51 AW583777.1		4507798 NT	AA233352.1	A1492415.1	2.0E-51 AW137826.1	AI732851.1	A1732854 4	BE782015.1	AF219927.1
	Most Similar (Top) Hit BLAST E Value	6.0E-51	6.0E-51	6.0E-51	6.0E-51	5.0E-51								3.0E-51	3.0E-51	3.05-54	3.0E-51	3.0E-51	L	Z.0E51	2.0E-51	2.0E-51	2.05-51	2.05-51		2.0E-51	2.0E-51
	Expression Signal	0.67	1.51	1.52	1.52	11.81	1.96	0.95	0.99	8.67	1.52	1.52	4.18	3.65	1.97	6	4.68	0.47	700	2.01	5.16	1.57	0.76	0.7	ć	3.86	0.61
	S O		37169	37425		26207				28052			37203	26571	29659	33246				08/07	27124	29107	28820	30837	30838	31663	
	Exan SEQ ID NO:		23884	24115		13546		_ !		15315			23911	13908	17031		21430	25430		8		16469		18155	}	1	19894
	Probe SEQ ID NQ:	9662	11221	11515	11515	774	785	970	1603	2601	3925	3925	11249	1163	4292	7479	8738	9968	, c	200	1683	3716	4458	5352	5352	5925	7209

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- 1		┰	т	_	_	_	_	_	_	_															
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens cell recognition molecule Casar 2 (KIAAnaga) mRNA	601676787F1 NIH MGC 21 Homo septems CINA close 1846 CE 3047843 E	601676787F1 NIH MGC 21 Home septems cDNA close IMACE 3050413 5	Homo saplens disrupted in schizophrenia 1 (DISC1), mRNA	ts74a07x1 NCI_CGAP_GC6 Home saplens cDNA clone IMAGE:2236980 3' similar to SW:TRKC_HUMAN Q16288 NT.3 GROWTH FACTOR RECEPTOR PDECLIBSOD	MR3-HT0487-150200-113-001 HT0487 Homo sanions - DNA	Homo saplens mRNA for KIAA0457 profelin partial cde	AV682474 GKB Home saplens cDNA clone GKBAGER 5	EST91296 Synovial sarcoma Homo sapiems cDNA 5' end	0534f09.x5 NCI_CGAP_KId5 Homo sepiens cDNA clone IMAGE:1326609 3' similar to SW:NME1_MOUSE P35436 GLUTAMATE INMDA1RECEPTOR SUBI INIT FERSION 1 PRECIDED.	ob3409.x5 NCI_CGAP_KId5 Home sapiens cDNA done IMAGE:1326909 3' SIMIENT to SW:NME1_MOUSE P35436 GLUTAMATE INMDA! RECEPTOR SUBUNIT EPSILON 1 PRECI IRSOR.	Homo sapiens myeloid/lymphoid or mked-lineage leukemia (trithorex (Drosophila) homolog); translocated to, 4 (MLT4), mRNA	Homo sablens eukenontic translation initiation factor 46 (1.06-1.1444). Days	AV742248 CB Homo sanians cDNA clone CRERCO12 8	801464995F1 NIH MGC 67 Home saniens cDNA clone IMAGE 3889248 KI	b12056t Testis 1 Homo saplens cDNA clone b12058	te39g02.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IWAGE:2089108 31	7096b02.x1 NCI_CGAP_Ov18 Homo saplens cDNA clone IMAGE:3644091 3' cimilar to TR:P87892 P87892 PROTEASE	AV760590 MDS Hamo saplens cDNA clone MDSCBB02 6'	Ng10h04.r1 Soares fetal liver spleen 1NFLS Homo saplens oDNA clone IMAGE:196667 6' similar to SP:YGAF_ECOLI P37339 HYPOTHETICAL PROTEIN IN GABP 3'REGION .	0410h04.r1 Soares fetal Ilver spleen 1NFLS Homo sepiens cDNA clone IMAGE:196567 6' similar to SP:YGAF_ECOLI P37339 HYPOTHETICAL PROTEIN IN GARP 3'REGION ·	295507.s1 Soares, feta_liver_spleen_1NFLS_S1 Homo sepiens cDNA clone IMAGE:448500 3' similar to contains THR.t3 THR repetitive element:	rw21g02.s1 NCI_CGAP_GCB0 Homo sapiens cDNA clone IMAGE:1241138 3' similar to contains THR.ts THR repetitive element:	H.sapiens mRNA for laminin-5, alpha3b chain
igle Exon Pro	Top Hit Database Source	N-	EST HUMAN	EST HUMAN	N	EST HUMAN	EST HUMAN	Į.	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	F	L L	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN		N
ה 	Top Hit Acession No.	7662349 NT	-61 BE901994.1	-51 BE901894.1	11037064 NT	-61 Al917078.1	BE165980.1	2.0E-51 AB007926.1	-51 AV682474.1	-51 AA378559.1	-51 AI732851.1	-51 AI732851.1	11419159 NT	4503528 NT	1.0E-61 AV742248.1	1.0E-51 BE779039.1	F18862.1	-61 AI572532.1	1.0E-51 BF434359.1	-51 AV760590.1	52 R91638.1	52 R91638.1	-52 AA777621.1	1.	
	Most Similar (Top) Hit BLAST E Value	2.0E-51	2.0E-61	2.0E-51	2.0E-51	2.0E-61	2.0E-51	2.0E-51	2.0E-51	2.0E-51	2.0E-51	2.0E-51	2.0E-51	1.06-51	1.0E-61	1.0E-51	1.0E-51 T18862.1	1.0E-51	1.0E-51	1.0E-51 /	9.0E-52 F	9.0E-52 F	9.0E-52 A	8.0E-52	8.0E-52
	Expression Signal	1.06	1.72	1.72	96'0	1.45	5.68	9.0	1.77	2.67	8.52	8.52	2.1	5.74	20.32	1.52	4.1	0.94	0.81	. 2	1.39	1.39	5.36	66.6	1.65
	ORF SEQ ID NO:				34766	İ	35336	35355	36199	36232	30837	30838	31017	25571		30294	30769	33322	33619		36520	36621		25605	26915
	Exan SEQ ID NO:	20038				22090	22156	22172	22979	23016	18165	18155	24870	12934	14226	17684	18110	20219	20498	25434	23282	23282	24726	12963	14229
	Probe SEQ ID NO:	7357	8599	8288	8632	9412	9503	9619	10332	10370	11298	11288	12523	112	1478	4929	5305	7549	7803	1783	10587	10587	12301	84	1482

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Single Exon Probes Expressed in Brain

	NA NA	¥	≰	₹				1		1	1				l						Н		lÌ	[]
Top Hit Descriptor	Homo sapiens hypothetical protein FLJ13556 similar to N-myc downstream regulated 3 (FLJ13556), mRNA	Homo sapiens hypothetical protein FLJ13556 similar to N-myc downstream regulated 3 (FLJ13556), mRNA	Homo sapiens hypothetical protein FLJ13556 similar to N-myc downstream regulated 3 (FLJ13556), mRNA	Homo saplens hypothetical protein PLJ13556 similar to N-myc downstream regulated 3 (FLJ13556), mRNA	Homo sapiens transforming growth factor, beta-induced, 68kD (TGFBI), mRNA	Homo sapiens transforming growth factor, beta-induced, 68kD (TGFBI), mRNA	2059a06.r1 Soares_parathyrold_tumor_NbHPA Homo sapiens cDNA clone IMAGE:326678 5' similar to contains Alu repetitive element	QV3-BT0537-271299-049-007 BT0537 Homo saplens cDNA	Homo sapiens S184 gene, partial cds, PS1 and hypothetical protein genes, complete cds, and S171 gene,	gg44f04.x1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1838047.3'	1246h04./1 NCI_CGAP_Bm52 Homo septens cDNA done IMAGE:2231671 5' similar to SW:PGBM_MOUSE 0.05793 BASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN PRECURSOR;	Homo sapiens FSHD region gene 1 (FRG1), mRNA	Homo sapiens adenylate cyclase activating polypeptide 1 (pituitary) receptor type i (ADCYAP1R1) mRNA	Homo sapiens nucleoporin 155kD (NUP155) mRNA	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens phosphoribosy pyrophosphate synthetase associated protein 2 (PRPSAP2) mRNA	Homo sapiens phosphoribosyl pyrophosphate synthetase associated protein 2 (PRPSAP2) mRNA	601440687F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3915836 5'	Homo sapiens hydroxysteroid (17-beta) dehydrogenase 4 (HSD17B4), mRNA	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	Homo sapiens DNA for Human P2XM, complete cds	Homo saplens hypothetical protein FLJ10675 (FLJ10675), mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	Human endogenous retroviral DNA (4-1), complete retroviral segment
Top Hit Database Source	NT	NT	NT	LN	ΙN	IN	NAMIH TRA	EST HUMAN	TIV	EST HUMAN	EST_HUMAN	IN.	LN	TN	NT	IN	TN	EST HUMAN	NT	TN	L	LN	LN	TN
Top Hit Acession No.	11968028 NT	11968028 NT	11968028 NT	11968028 NT	11416585 NT	11416585 NT	W56471 1	6.0E-52 BE072409.1	A E400007 4	6.0E-52 AI208794.1	BE048172.1	-52 11437365 NT	4501922 NT	4758843 NT	4507500 NT	4506132 NT	4506132 NT	4.0E-52 BE622032.1	11417035 NT	11418177 NT	52 AB002059.1	11437042 NT	52 M10976.1	52 M10976.1
Most Similar (Top) Hit BLAST E Value	8.0E-52	8.0E-52	8.0E-52	8.0E-52	8.0E-52	8.0E-52	7 OF-82	6.0E-52	A 70 A	6.0E-52	6.0E-52	5.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	4.0E-52	3.0E-52	2.0E-52	2.0E-52
Expression Signal	3.13	3.13	9.9	6.6	0.67	0.67	20 %	0.78	7.07	0.86	28.	9.0	1.32	1.02	0.99	1.33	1.33	1.74	5.48	5.11	5.23	10.57	2.88	2.88
ORF SEQ ID NO:	27085	27086	27085	27086		33179	34745		027430		37119		27165		29297				34263					25963
Exon SEQ ID NO:	14396	14398	14396	14396	20094	20094	21602	13918	l	1	23837	21959	14466	14521	16656	18012	18012	20833	21125	24631	24930	16815	13332	13332
Probe SEQ ID NO:	1650	1650	3976	3976	7417	7417	8944	1184	7	5641	11170	9292	1723	1780	3906	5204	5204	7938	8432	12143	12627	4071	549	549

Page 308 of 536 Table 4 Single Exon Probes Expressed

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	bb66b07.y1 NIH_MGC_9 Homo sapiens oDNA clone IMAGE:3030421 6' similar to ab:X18483 M muschilling	mrana for 2pt-1 zinc finger protein (MOUSE),	002084/10F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4248891 6	Nove human gene mapping to chromosome 20, similar to membrane transporters	Hacoedo.sr Scares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1690784 3'	gaodeus, st Soares NhHMPu_S1 Homo saplens cDNA clone IMAGE:1690784 3'	ICCOLOGIFTS IN SECTION STATEMENT OF THE SECTION OF	Homo sapiens interieutin 21 receptor (IL21R), mRNA	como saprens mixina for Kilaa 1081 protein, partial cds	Maccondition Court Briz Homo sapiens cDNA clone IMAGE:1608311 6	wacaca muana bera-ubulin mRNA, complete cds	47-300.5.1 States, retal, liver_spieen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:463272.3. Them capiens NADH dehydrogenase (ubiquinone) Fe-S protein 5 (15kD) (NADH-coonzyme Q reductions).	(WOUTSS) MANA	normo septems SE I domain and mariner transposase fusion gene (SETMAR) mRNA	Promo septens SE I domain and mariner transposase fusion gene (SETMAR) mRNA W49c04.x1 NCL_CGAP_Lu19 Homo septens cDNA clone IMAGE:2408150.3' similar in contains TLIB to	THR repetitive element;	WyseCV4.X1 NCI_CGAP_Lu19 Home sapiens cDNA clone IMAGE:2406150 3' similar to contains THR.b2	AV715377 DOB Home septem contact in the september of	449012 11 Spars febiling NEULINOS INCOME UCBAIEUS 3	Homo saplens LIM domain kinase 2 (11MK2) mBNA	xn72e07.x1 NCI_CGAP_CML1 Home sapers CONA clone IMAGE:2700036 3' similar to contains Atu	wf67d05.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE.2360649 3' similar to TR:O18889	GIOSOS CARBOXYLES ERASE;	All 3112.81 Sogies, testis, NHT Homo saplens cDNA clone IMAGE:743879 3'	Homo septens glutamate-animonia ligase (glutamine synthese) (GLUL) mRNA	politico captario al yeuratasso D (ARSD), transcript varient 1, mRNA politico estanscriptase homolog (retroviral element) (numan, endocenous retroviral element 1974), Una	Genomic, 660 ntj	Fruman Prepiscoprotein (MDR1) gene, exon 4
gle Exon Pro	Top Hit Database Source	TO D	EST TOWAN	FIGURAIN HA	EST LIBARE	Т	Т	AIUMOI -		HIMAN	Т	HIMAN					EST_HUMAN	EST HUMAN 17	Т	Г		EST_HUMAN R		1	NON-				
Sin	Top Hit Acession No.	.50 RF 207474 4	T	T	T	T	Ţ	1868	52 AB029004 1	1		-52 AA778795.1	2780	5730038 NIT	TIMBEOORE						17990	2.0E-52 AW236297.1 E		T	acore	4502238 NT	0.000		U38964 1
	Most Similar (Top) Hit BLAST E Value	2.05.52	2 0F-52	2.0E-52	2.0E-52	2.0E-52	2.0E-52/	2.0E-52	2.0E-52	2.0E-52	2.0E-52 A	2.0E-52	2.0E-52	2 0E-52	2.0E-52	2 OF-52 Aleaster 4	4.0E-32	2.0E-52 A	2.0E-52 AV715377.1	2.0E-52 W	2.0E-52 114	2.0E-52 A\	2.0E-52 AIB08985 1	1.0E-52 AA63445 1	1.0E-52	1.0E-52	4 0 1 10 6	1.0E-52 M2	1.0E-52 U3
	Expression Signal	2.04	6.03	2.13	1.29	1.28	4.11	1.86	0.99	1.17	9.03	0.84	0.88	5.53	5.53	3.45		3.15	3.09	1.72	2.76	24.36	4.49	1.96	37.84	6.0	2.87	4.35	2.51
	ORF SEQ ID NO:	27963		30260		30288	31326	32024	32390	32571		34672		35879	35880	37111		37112	37131			30601		25937	26773		28465	30684	32059
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	Probe SEQ ID NO:	2503	2740	4920	4952	4952	5617	6274	6613	8843	8651	8834	9379	10015	10015	11165		11165	111/8	222	17678	11981	12350	520	1350	2537	3055	5250	6300

Page 309 of 536 Table 4 Single Exon Probes Expressed in Brain

Probe Exon SEQ ID SEQ I

Page 310 of 536 Table 4 Single Exon Probes Expressed in Brain

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Cingre Lyon Probes Expressed in Brain	Top Hit Descriptor	FST77595 Pancians times III U	Homo saplens Button's fuscine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein [144] and FTP3 (FTP3).	Company of the second second second (vacuolar proton pump) 31kD; Vacuolar proton-ATPase, submit E-VATPase and company 1 proton pump) 31kD; Vacuolar proton-ATPase, submit E-VATPase	Homos explains core-binding factor, runt domain, alpha subunit 2; translocated to, 1; cyclin D-related	Home spiens core-binding factor, runt domain, alpha subunit 2; translocated to, 1; cyclin D-related	Home action of the state of the	Himen Kniesen Files - Bass 11. 1.	PM4. CTrage 470000 and 200 officer (1F34) gene, partial cds	PM4-CTAsse-170800 004 202 CT0386 Homo sapiens cDNA	Fig. 1 constant in control i gus C i Used Homo sapiens cDNA	2822885 Entime NILL MCC 7 1	Pomo sociolo No. Mo. / Home sapiens cDNA clone IMAGE:2822665 5'	Homo saptens Ad pseudoautosomal region; segment 2/2 Homo saptens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes.	CALL VILLAGE CAS	CM HAIN 1028-150800-543-602 NN1029 Homo sapiens cDNA	INSTACTOR INSERTINGUEST-DUT BIN1058 Homo sapiens cDNA	Habriers PNA for habrier	H. Saplens mRNA for mysin. IF	H. sapiens mRNA for myosin-IF	2822943.3prime NIH MGC 7 Home sanieps cDNA close IMA CE:382324 al	Homo saplens (O motif containing GTDs as equinities as et al. (1000 cm.)	601272863F1 NIH MGC 20 Home septeme abala Committee Management (ICCAP1) mRNA	Homo sapiens insulin-like growth factor 2 recenter (10 Eng) - Pata	Homo sapiens ATP-binding cassette, sub-family A (ABCI)	al78c12.s1 Soares_testis_NHT Homo sapiens cDNA clone 1377046 3' similar to contains MER30.t3 MER30	Homo seniens mRNA for monorite chemicatories and in the control of	W68412.s1 Society placents at 1909 events 2NbHP8to9W Home sapiens cDNA clone IMAGE:257399.31	יייי יפור יפור יפור יפור יפור יפור יפור
שונה באחוו בוס	Top Hit Database Source	EST HUMAN	- LN	L L	LN	F	FZ	L	T HIMANI	Т	Т	Т	Т		1	IN TOWAR	┰				EST_HUMAN		T HUMAN	9		EST HUMAN	Т	HOMAN	7
5	Top Hit Acession No.	-53 AA36656.1	2.0E-53 U78027,1	4502316 NT			AF08382	T	Ţ	T		Γ	1.0E-53 AJ271736 1		-53 RF384204 4	T	T	T		Γ	53 AW245422.1 E	4506786 NT	54 BE386785.1 E	34610	6005700 NT	7.0E-54 AA812637.1 E		74 N27177.1 E	
	Most Similar (Top) Hit BLAST E Value	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	2.0E-53	1.0E-53	1.0E-53	1 OF 53	1,0E-53	1.0E-53	1.0E-53)	1.0E-53)	1.0E-53	1.0E-53 /	9.0E-54	8.0E-54	8.0E-54	8.0E-54	7.0E-54 A	7.0E-54 Y	7.0E-54 N	
	Expression Signal	5.82	2.78	8.73	1.48	1.46	3.72	2.63	2.67	2.67	-	3.82	1.51	1.08	1.62	0.68	0.54	5.91	1.41	1.41	2.29	6.16	2.4	1.77	26.87	1.27	1.65	6.38	
	ORF SEQ ID NO:		27788		28172	28173	28851	29411	30804	30805	33590		26860	28813	32370	32903	33658	34826	37757	37768	37255	30651	25655	27278	31588	25845	27273	27867	
	S	13231	15052	15253	15438				18143	18143	20468	21975	14175	16162	19356	19834	20531	21677	24417	24417	24507	300	13015	14568	18633	13200	14581	14930	•
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Page 311 of 536 Table 4 Single Exon Probes Expressed in

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| Top Hit Descriptor | Homo sapiens similar to nuclear factor related to bonno R binding systems (1) Account (2) | Homo sapiens goldin-like protein (GLP) mRNA | Homo sapiens golgin-like protein (GLP), mRNA | qb67g03.x1 Soares fetal_heart_NbHH19W Homo septens cDNA clone IMAGE:1705204.3' similar to contains OFR.tt OFR repetitive element: | Homo sapiens jun dimerization protein gene, partial cds; cfos gene, complete cds; and unknown gene | Homo sapiens iun dimentzation brotein dene nartial cds. ofte dene commiste ode, and university | Homo sapiens DNA for MICB, exon 4, 5 and partial cds | Homo sapiens hypothetical protein DKFZnd24Mn35 (DKFZnd24Mn35) mBNA | Homo sapiens hypothetical protein DKFZp434M035 (DKFZp434M035) mRNA
 | Homo saplens hypothetical protein DK FZp434M035 (DKFZp434M035), mRNA | Homo sapiens chloride channel 6 (CLCN6) mRNA | AV754746 TP Homo sapiens cDNA clone TPGAAC10 5 | Homo sapiens phosphatidylinositol 4-kinase, catalylic, alpha polymentida (PIKACA) mBNA | H. sapiens sho pseudocene, p66 isaform | RC3-ST0197-151099-011-f08 ST0197 Homo sapiens cDNA | ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2)

 | Tupala belangeri beta-actin mRNA, partial cds | EST177696 Jurkat T-cells VI Homo sapiens cDNA 5' end similar to giyceraldehyde-3-phosphate | dehydrogenase | Human mb/s & Visabut/ gene, parter das | Halled Hill of Science NET T CBC St. 1

 | MOZZI FINE SOME STATE JOHN STATE TOTAL SEPTENS CONA CIONE IMAGE: 2329269 3' Similar to TR: 002711
 002711 PRO-POL-DUTPASE POLYPROTEIN : | EST185371 Colon carcinoma (HCC) cell line Homo saplens cDNA 5' and | hd87g08.x1 NCL CGAP GC6 Homo saplens cDNA clone IMAGE: 2016542.3. | DKFZp434E0731 r1 434 (synonym: htts3) Homo septiens cDNA clara DKFZzd424E0734 g | IL-8T189-190399-007 BT189 Homo saniens cDNA | Homo sapiens BMX non-receptor tyrosine kinase (BMX) mRNA
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 | 0.77 | 1.91 | 0.86 | 1.78 | 1.15 | 1.51 | 3.78

 | 13.34 | | 3.72 | 322 |

 | - | 4.47 | 0.91 | 0.96 | 1.34 | 1.74
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 | 28688 | 29368 | 29794 | 30145 | | 36430 | 27611

 | | 00000 | 27.748 | 27249 |

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 | 16038 | 16734 | 17165 | 17523 | 17550 | 23199 | 14876

 | 12890 | 42.700 | 14538 | 14538 |

 | 15962 | 12918 | 14312 | 15288 | 15342 | 18603
 | 19971 | 19971 |
| Probe
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 | 3277 | 3986 | 4429 | 4792 | 4819 | 11432 | 2146

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Page 312 of 536 Table 4 Single Exon Probes Expressed in Brain

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	Top Hit Descriptor	Homo saplens golgi autoantigen, golgin subfamily a 6 (GOI GAR) mRNA	802019408F1 NCI CGAP Bm67 Homo septens cDNA clone IMAGR-4455121 5	270f12.r1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:727727 6' similar to TR:0191315 0191315 ANDROGEN.DEPENDENT EXPRESSED PROTEIN	EST388829 MAGE resembles MAGO Liver contract CNA	RC1-BT0313-131199-011-b08 BT0313 Homo segiens colva	Homo sapiens killer cell lecthalike recentor subfamily C. mamber 1 (v. 1504)	Homo saplens nuclear antiden So 100 (SP100) mRNA	nt78a09.s1 NCI_CGAP_Pr3 Homo sepiens cDNA clone IMAGE:1204600 similar to contains element L1 repetitive element :	ลย22g03 y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783764 5' similar to SW:CUL1 HUMAN 013616 Cult tin HOMoi กิด 1	Homo saplens chromosome 21 segment HS21C010	wy60b12.x1 Soares_NSF_F8_9W_OT_PA_F31 Homo sapiens cDNA clone IMAGE:2652927 3' similar to TR:082084 Q62084 PHOSPHOLIPASF C NEIGHBORING	Homo sapiens mRNA for phospholipase C-beta-16 (PI CR1 pena)	nj45g09.s1 NCI_CGAP_Pr9 Homo sapiens cDNA clone IMAGE:895488 similar to gb:X53777 60S RIBOSOMAL PROTEIN L23 (HUMAN):	Homo saplens chaperonin containing T-complex subunit 6 (CCTR) mRNA	Homo saplens peptidylarginine deiminasa hone III (1 OC\$1702) mRNA	Homo saplens small Inducible cytokine subfamily A (Cys-Cys) member 14 (SCYA14) mBNA	243c11.y1 NCL CGAP_Brn52 Homo saplens cDNA clone IMAGE:2291348 6'	Homo sapiens KIAA0100 gene product (KIAA0100), mRNA	Homo sapiens mRNA for KIAA1591 protein, partial cds	Homo sapiens mRNA for KIAA1591 protein, partial cds	Homo sapiens EVI5 homolog mRNA, complete cds	Homo saplens mRNA for KIAA0995 protein, partial cds	Homo saplens mRNA for KIAA0995 protein, partial cds	Homo saplens neurofibromin 1 (neurofibromatosis, von Recklinghausen disease, Watson disease) (NF1).	mRNA	Homo saplens mRNA for brain ryanodine receptor, complete cds	Homo saplens Janus kinase 2 (a protein tyrosine kinase) (JAK2), mRNA	Hamo saplens serologically defined colon cancer antigen 10 (SDCCAG10), mRNA	Homo saplens serologically defined colon cancer antigen 10 (SDCCAG10), mRNA
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	Expression Signal	1.63	4.93	3.26	2.98	2.51	8.86	0.98	1.37	1.22	1.85	1.52	1.18	3.2	2.08	1.02	1.64	1.2	5.04	13.99	13.99	0.68	0.65	0.65		8.8	4.11	0.79	1.01	1.01
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	Probe SEQ ID NO:	10949	11024	11341	12056	12097	827	1344	1539	2541	2608	2896	3311	3536	4181	4826	5388	5518	6976	6771	5771	8229	6713	6713	000	207	9529	808	10021	10021

Page 313 of 536 Table 4 Single Exon Probes Expressed in Brain

Page 314 of 536 Table 4

Single Exon Probes Expressed in Brain	Top Hit Descriptor		Home sapiens mRNA for KIAA0611 protein, partial ods	Homo saplens nel (chicken)-like 2 (NELL2), mRNA	Homo saplens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA	Homo sapiens RNA binding modif protein, Y chromosome, family 1, member 41 (PRAZVA 41) - BAIA	Homo saplens predicted osteoblast protein (GS3786), mRNA	Homo sepiens predicted ostaobiast protein (GS3786), mRNA 7/5/25/10/X/ Soares NSF FR ow OT DA P SA LI	contains L1, 13 L1 repetitive element.	Homo saplens proteasome (prosome, macropain) subunit alpho to 2 (DEXIAA).	Homo sapiens proteasome (prosome, macropain) subjuit chest.	Homo sapiens diacyglycerol kinase, damma (90kh) (10kh) (10kh)	Homo saplens diacylglycerol kinase, gamma (90km) (100kg) mpNA	Homo saplens ubiquitin-conjugating enzyme E2 variant 1 (1 IBE 2014) - BN A	Homo saplens Xq pseudoautosomal region segment 1/2	Homo saplens chromosome 21 serment H231/CA1	43c5 Human retina cDNA randomly primad autilibrary Home continued to the continue of the conti	601886575F2 NIH MGC_17 Homo sapiens cDNA clone IMAGE:4132222 F	PM1-HT0603-090300-001-408 HT0663 Home serious CDNA	Homo sapiens chromosome 21 segment HS21C084	Human endogenous retrovirus pHE.1 (ERV9)	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo sapiens syntadn-binding protein 1 (STXBP1) mRNA, and translated products	CM1-H 10876-150800-357-g03 HT0876 Homo sapiens cDNA	UI-TIT-DNU-aKS-1-08-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3078275 5	hrzahos v Nici CCAP Kid11 Homo sapiens cDNA clone IMAGE:3134463 3'	am98h05.s1 Strategene schizo brain S11 Homo saplens cDNA clane IMAGE:3134463 3'	THR.b2 THR repetitive element;	QV0-BN0147-280400-213-g06 BN0147 Homo saniens CONA	AU119344 HEMBA1 Homo sepiens cDNA clare HEMBASSES ::	Homo saplens mannose-6-phosphate receptor (cation dependent) (Mebb) ENA	WALKER (V. 1944) (V. 1944)	Prychagus Cuniculus New Zealand white elongation factor 1 alpha (Rabella2) mRNA, complete cds Homo saplens mRNA for KIAA0903 protein, partial cds
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	ORF SEQ ID NO:		35984		26075		26854		207720	27482	27542	27542	27.7.6							25807		26049	30080	33162	34804	34805			38708	8 22 23	90007	25636	26543
	SEQ ID NO:	7 22585	L	L	13434	L	Ц	14250		14754	L	1			1			24538	24003	13164	13321	13413	17455	25113	21664	21664	21747	21828	23550	12024	707	12997	13883
	Probe SEQ ID NO:	8937	10122	12137	927	1421	1421	1504	2040	2019	2070	2078	2308	2698	8242	146	12057	11998	12721	368	838	634	4723	7403	8963	8863	8906	9140	10870	90		184	1127

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplane bate this ille m DNA consistent	6018R204GET NIH MOC ES LICE COLLEGE CAS	Homo saplens in thulin hate polymentide (Tillips) Table	Homo saplens tribulin, beta polypeptide (TUBB) mRNA	Homo saplens X-linked anhidrolito ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat regions	Wb08f08.x1 NCI_CGAP_GC8 Home sapiens cDNA clone IMAGE:2306191 3' similar to SW:DCOR_MUSPA P27119 ORNITHINE DECARBOXM ASE ·	WD08f08.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2305191 3' similar to SW:DCOR_MUSPA P27149 ORNITHINE DECARBOXX ASE.	Homo sablens (incharacted tool box morre:	Homo Saplens Incherecterized beneating the protection of the complete cds	Homo carlens hambourds account and a figure of a figur	m85012 v1 NCL CGAD Brose Lorent 1 (LSP1) gene, LSP1-7 allele, partial cds	maga12x1 NCI CGAP Brack Home contact that a little contact to the	Homo sablens hypothetical profels DBO4304 (DBO4304) - DBA4	EST28889 Caraballum II Homo souless apply #1 and 1844, mrnA	EST28889 Carabellum II Homo septem ADNA 51 and	Homo sanians MHC class 1 varions	Homo sepiens Down syndrome candidate realize 7 (DSCB4) DNA	Homo sapiens chromosome 21 segment HS21 Ches	Homo sapiens suberkiller viralicidic activity 2 (S. cerawisiae homologo, lika (SK)) voi 1	Homo sapiens sparc/ostbonectin, cwov and kazal-like domains anyward (was transfer or to construct or to constr	MKNA MENON (PSSUCER) (SPSUCER) (SPSUCER) (SPSUCER)	Home suprems sparciosceneral, owov and kazal-like domains proteoglycan (testican) (SPOCK) mRNA	Homo septems 1) social metarge octated membrane protein 2 (LAMP2), mRNA Homo septems I IM hindian domain 2 /I DB21. —DNA	Homo sapiens LIM binding domain 2 (LDB2) mRNA	Homo saplens bone morphogenetic profein 5 (RMDs) mRNA	Homo saplens mRNA for KIAA0145 protein partial cds	Homo sapiens KIAA0317 gene product (KIAA0317), mRNA	Homo sapiens mRNA, similar to rat myomegalin, complete ods	Homo saplens nuclear pore complex interacting protein (NPIP), mRNA
gle Exon Prob	Top Hit Database Source	TN	T HI IMAN			۲	EST HUMAN	Ì	Т			┰	EST HUMAN		HIMAN	Т	Т										E L			
Sin	Top Hit Acession No.	56 AF141349 1		077728	4507728 NT	56 AF003528.1	56 A1632488.1	56 A1632488.1	Ì	-56 AF217508.1		l		24029	3.0E-56 AA325826.1			37042	3.0E-56 AL163268.2	5902085 NT	4759163 NT	1750183 NT	11421124 NT	4504970 NT	4504970 NT	11418704 NT		11434956 NT	3042556.1 NT	5902013 NT
	Most Similar (Top) Hit BLAST E Value	4.0E-56	4.0E-56	4.0E-58	4.0E-56	4.0E-56	4.0E-56	4.0E-56	4.0E-56	4.0E-56	4.0E-56/A	4.0E-56 A	4.0E-56 A	3.0E-56	3.0E-56 A	3.0E-56	3.0E-56 A	3.0E-56	3.0E-56 A	3.0E-56	3.0E-56	3.05-58	3 0E-56	3.0E-56	3.0E-56	3.0E-58	3.0E-56 D63479.2	3.0E-56	3.0E-56 AB042556.1	3.0E-56
	Expression Signal	8.58	2.69	7.28	7.28	3.49	1.48	1.48	6.01	6.01	2.02	8.88	8.88	4.17	1.54	1.54	1.61	1.43	4.27	2.34	2.12	212	7.03	1.15	1.15	4.68	0.85	1.38	1.71	6.37
	ORF SEQ ID NO:	25470		28157	28158	25929	28074	28075	31912	31913	36266	36764	36765	26742	28527	28528	-	29728	29759	29902	31302	31303	32547	32981	32982	34550	35562	36240	36571	37239
	SEQ ID NO:	12854	14902	15419	15419	13297	15331		18941	18941	23049	23523	23523	14068	15887	15887	16587	17093	17127	17269	18383	18383	19519	19908	19908	21407	22364	23026	23333	23840
	Probe SEQ ID NO:	26	2173	2712	2712	2815	2836	2836	6164	6164	10403	10841	10841	1319	3122	3122	3815	4355	85 865	4534	5598	5598	6775	7223	7223	8715	9713	10379	10642	11284

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Top Hit Descriptor		Homo saplens nuclear pore complex interacting protein (NPIP), mRNA	Homo sapiens NACP/alpha-synuclein gene, exon 5	Homo sapiens NACP/alpha-synuclein gene, exon 5	Homo sapiens caveolin 3 (CAV3), mRNA	Homo sapiens caveolin 3 (CAV3), mRNA	T		Г	Homo sapiens mRNA for KIAA1414 protein, partial cds	Г		Macaca fascicularis protein tyrosine phosphatase (PRL-1) mRNA, complete cds	Г	Г	I QV-8T077-130199-079 BT077 Hamo saplens cDNA	Γ	┱	Т	I RC2-CT0163-220999-001-E02 CT0163 Hamo sepiens cDNA		Homo sapiens EphA4 (EPHA4) mRNA	Homo saplens EphA4 (EPHA4) mRNA	Homo sapiens serine protease 17 (KLK4) gene, complete cds	Homo sapiens serine protease 17 (KLK4) gene, complete cds	Home sapiens mRNA for cyclin B2, complete cds	QV4-ST0234-181199-037-f05 ST0234 Homo sapiens cDNA		Т	╗	Homo sapiens EphA4 (EPHA4) mRNA	Homo sapiens EphA4 (EPHA4) mRNA	Homo sapiens glutamate receptor, Ionotrophic, AMPA 4 (GRIA4) mRNA	Homo saplens aconitase 2, mitochondrial (ACO2), mRNA	Homo saplens mRNA for KIAA0898 protein, partial cds
Top Hit Database Source		Ā	ΤN	L'N	Į.	N-	EST_HUMAN	EST_HUMAN	EST HUMAN	Ľ	EST_HUMAN	N TN	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	N FN	EST HUMAN	FN	EST_HUMAN	EST_HUMAN	TN	TN	L	TN	NT	EST_HUMAN	MANUAL TOO	אאוסט ביים	EST_HUMAN	칟	NT	ΤN	NT	TN
Top Hit Acession No.		5902013 NT	56 U46900.1	56 U46900.1	11434876 NT	11434876 NT	56 AA199818.1	56 BE064386.1	56 BE064386.1	56 AB037835.1	56 AV703184.1	5730038 NT	AF190930.1	-56 AW589833.1	56 AW589833.1	36 Al905162.1	6681002 NT	56 AW609520.1	1.0E-56 AL163203.2	1.0E-56 AW845987.1	57 AW880885.1	4758279	4758279 NT				8.0E-57 AW816405.1	AW(264500 4	O'OE-SI AW ZOTSES.	AA49610		4758279 NT	4557630 NT	8.0E-57 11418185 NT	П
Most Similar (Top) Hit BĽAST E	Value	3.0E-56	3.0E-56	3.0E-56	3.0E-56	3.0E-58	2.0E-	2.0E-56	2.0E-56	2,0E-56	2.0E-58	2.0E-56	1.0E-56	1.0E-56	1.0E-56	1.0E-56	1.0E-56	1.0E-56	1.0E-56	1.0E-56	9.0E-57	9.0E-57	9.0E-57	9.0E-57	9.0E-57	9.0E-57	8.0E-57	D 0	0.000	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57
Expression Signal		6.37	1.74	1.74	1.52	1.52	1.7	1.05	1.05	1.18	1.64	1.47	1.84	2.15	2.15	0.99	0.97	0.57	0.59	1.71	2.52	1.14	1.14	2.17	2.17	1.48	3.01	86.8	200	1.51	0.98	0.98	1.3	3.29	1.85
ORF SEQ ID NO:		37240		37591	31083	31084					28934	32731		29056	29057	30303	30453	32589		35800		29548	29549	37134	37135	37420	25738	26303	20002	2/204	28791	28792	30205	30728	32083
Exen SEO ID NO:		23945		24268						15753				16417	16417	17696	17836	19558	22505	22596	13389	16920	16920	23848	23848	24107	13096	12823	335	-40 -04 -04 -04 -04 -04 -04 -04 -04 -04	16135	16135	17682	25276	19078
Probe SEQ ID NO:		11284	11673	11673	12095	12095	511	716	716	2987	3523	0669	929	3664	3664	4972	5118	6724	9855	9948	611	4180	4180	11183	11183	11506	280	788	3 8	ang.	3376	3376	4852	5161	9306

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens mRNA for KIAA0960 protein, partial cds	Homo sapiens mRNA for KIAA0960 protein, partial cds	Homo saplens KIAA0718 gene product (KIAA0716), mRNA	Homo saplens mRNA for KIAA0837 protein, partial cds	Homo saplens mRNA for KIAA0837 protein, partial cds	Homo sapiens hypothetical protein FLJ20371 (FLJ20371), mRNA	Homo sapiens SH3-domain binding protein 1 (SH3BP1), mRNA	Homo sapiens SH3-domain binding protein 1 (SH3BP1), mRNA	Homo sapiens mRNA for KiAA1342 protein, partial cds	Homo saplens smg GDS-ASSOCIATED PROTEIN (SMAP), mRNA	Homo sapiens ang GDS-ASSOCIATED PROTEIN (SMAP), mRNA	Homo saplens NME7 (NME7), mRNA	Homo sapiens NME7 (NME7), mRNA	Homo sapiens Kruppel-like factor 8 (KLF8), mRNA	Homo sapiens phosphatidylinositol 4-kinase 230 (pl4K230) mRNA, complete cds	Homo saplens phosphatidylinositol 4-kinase 230 (pi4K230) mRNA, complete cds	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens large conductance calcium- and voltage-dependent potassium channel alpha subunit (MaxK)	Homo saplens Xq pseudoautosomal region; segment 1/2	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	Homo sapiens ubiquitin protein ligase E3A (human papilioma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA	no13f07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1008037 similar to SW:RS10_HUMAN P46783 40S RIBOSOMAL PROTEIN S10.:	EST64770 Hippocampus II Homo sapiens cDNA 6' end	733b10.x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:3296443 3' similar to WP:Y47H9C.2	CE20283 ;	7/33b10.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:3296443 3' similar to WP:Y47H9C.2 CE20263 ;	Homo sapiens cell-line tsA201a chloride ion current inducer protein ((CIn) gene, complete cds	RC3-CT0254-110300-027-410 CT0254 Homo saplens oDNA
le Exon Probe	Top Hit Database Source	Ĭ				Ĭ				H							H LN	Į.	Y E		子 S		EST HUMAN P4	Т		EST HUMAN CE	FST_HUMAN CE	NT Ho	EST_HUMAN RC
Sing	Top Hit Acession No.	57 AB023177.1	-57 AB023177.1	7662263 NT	8.0E-57 AB020644.1		8923349 NT	11545732 NT	11545732 NT	37 AB037763.1	7857592 NT	7657592 NT	7242158 NT	7242168 NT	TN 6265009	7.0E-57 AF012872.1		7.0E-57 AF020503.1		-		4507798 NT		3.0E-57 AA348335.1 E		3.0E-67 BE676622.1	3.0E-57 BE676622.1 E	3.0E-57 AF232708.1 N	П
	Most Similar (Top) Hit BLAST E Value	8.0E-67	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57	8.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57 U11058.2	6.0E-57 /	4.0E-57	3.0E-57	3.0E-57	3.0E-57 A	1000	3.0E-67 E	3.0E-57 B	3.0E-57	3.0E-67 A
	Expression Signal	12.87	12.87	0.64	1.7	1.7	3.29	2.74	1.69	2.07	1.71	1.71	6.0	6.0	1.08	1.39	1.39	0.95	0.95	2.63	1.67	0.78	16.24	2.99		0.95	0.95	1.74	62.34
	ORF SEQ ID NO:	32137	32138	33107	33423	33424	30487	31028	31028		28093	28094	28655	28656	28677	29246	29247		30089	-	29125	28220		27848	207	28151	28152	28955	
	Exon SEQ ID NO:					20312	17889	24828	24828	25060	16350	15350	16006	16006	16027	16608	16608	17135	17462	26310	16489	13568	14056	15111	4644	2	15414	16305	16438
	Probe SEQ ID NO:	6372	6372	7349	7848	7648	11460	12429	12473	12820	2639	2639	3244	3244	3265	3858	3858	4398	4730	12785	3736	786	1308	2390	2707))/2	2707	3550	3685

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Top Htt Descriptor	Homo sapiens anglotensin I converting enzyme (peptidyl-dipeptidase A) 2 (ACE2), mRNA	601569896F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3944302 5'	42f6 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA	Homo saplens hypothetical protein FLJ11656 (FLJ11656), mRNA	Homo sapiens hypothetical protein FLJ11656 (FLJ11656), mRNA	Homo sapiens KIAA0649 gene product (KIAA0649), mRNA	Human famesyl pyrophosphate synthetase mRNA, complete cds	AU117659 HEWBA1 Homo saplens cDNA clone HEMBA1001910 5'	Homo sapiens hypothetical protein FLJ11656 (FLJ11658), mRNA	Homo saplens hypothetical protein FLJ11656 (FLJ11658), mRNA	2820473.5prime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2820473 5	2b45d11.r1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:306549 5	RCO-HT0112-080999-001-C06 HT0112 Hamo sapiens cDNA	AJ003649 Selected chromosome 21 cDNA library Homo sapiens cDNA clone MPIpI10-1L1	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C004	MR0-BT0551-060300-103-b03 BT0551 Homo seplens cDNA	Homo sapiens chromosome 21 segment HS21C083	ze31c05.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:380584 5' similar to contains L1.t3 L1 receititive element :	7n80f04.x1 NCI_CGAP_Ov18 Homo saplens cDNA clone IMAGE:3570966 3' similar to contains TAR1.t1	WERZ repetitive element;	Homo sapiens small inducible cytokine subfamily A (Cys-Cys), member 22 (SCYA22), mRNA	Homo sapiens cell-line KG1 transcriptional regulatory protein p54 mRNA, complete cds	Homo saplens 17-beta-hydroxysteroid dehydrogenase IV (HSD17B4) gene, exons 3 and 4	Homo sapiens KIAA1065 protein (KIAA1065), mRNA	Homo sapiens KIAA1065 protein (KIAA1065), mRNA	Homo sapiens hypothetical protein FLJ20041 (FLJ20041), mRNA	Homo saplens hypothetical protein FLJ20041 (FLJ20041), mRNA	ho32a08.X1 NC_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3039062 3' similar to TR:O00246 O00246 HYPOTHETICAL 9.3 KD PROTEIN :	hs33d06.x1 NCI_CGAP_Kid12 Homo sepiens cDNA clone IMAGE:2875499 3' similar to contains THR.b3 THR repetitive element ;
Top Hit Database Source	LN	EST_HUMAN	EST_HUMAN	NT	Z	LN	N	EST_HUMAN	N	N	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	Z	. LN	NT	EST_HUMAN	TN	EST HUMAN		EST_HUMAN	NT	LΝ	NT	NT L	Z	NT	N N	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	11225608 NT	-57 BE796537.1	-57 W28130.1	11545798 NT	11545798 NT	11427757 NT	-57 J05262.1	57 AU117659.1	11545798	11545798 NT	3.0E-57 AW248374.1	3.0E-57 W23871.1	-57 AW178575.1	3.0E-57 AJ003649.1	2.0E-57 AF246219.1	-57 AF246219.1	-57 AL:163204.2	-57 BE073264.1	-57 AL163283.2	57 AA016131.1		-57 BF115266.1	11431281 NT	57 AF045452.1	57 AF057722.1	11434330 NT	11434330 NT	11424084 NT	11424084 NT	57 BE043031.1	57 AW470791.1
Most Similar (Top) Hit BLAST E Value	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	3.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57		2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	1.0E-57	
Expression Signal	1.24	3.23	3.77	2.18	2.16	0.7	0.73	4.17	1.03	1.03	2.85	8:38	2.32	1.48	1.39	1.39	1.24	0.79	6.73	184		33.81	0.66	1.08	1.86	0.49	0.49	2.42	2.42	3.62	6.08
ORF SEQ ID:NO:	31680	31773	33871	33892	33893		34157	34596	35047		36747	30616		31010	26919	26920		29301					31810	34363	35602	36338	36399	37189	37190		
Exon SEQ ID NO:	18721	18813	20738	20764	20764	ı	ı	21449	!	ı	23508		L	24928	14234	14234	16188	16660	17209	! .			18846	21221	22397	Ι.	ı	j	L	l	
Probe SEQ ID NO:	5939	6033	8044	8070	8070	8179	8328	8757	9149	9149	10825	12101	12460	12623	1487	1487	3432	3910	4474	5582		5943	2909	8529	9746	10525	10525	11238	11238	8593	12249

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Table 4
Single Exon Probes Expressed in Brain

		_	_				_	_			_	_	_																
	Top Hit Descriptor	EST11348 Uterus Hamo saplens cDNA 6 end	1801309465F1 NIH MGC 44 Homo septens cDNA rione IMAGE 3834000 F	601445948F1 NIH MGC 65 Hamo saplens cDNA clone IMAGE 3850211 5	t34b07.x1 NCI_CGAP_Ov23 Homo sapiens cDNA clone IMAGE:2220181 3' similar to TR:O16476 O16476 UNNAMED HERV-H PROTEN:	tr34b07.x1 NCI_CGAP_Ov23 Homo sepiens oDNA done IMAGE:2220181 3' similar to TR:016476 O16476 UNNAMED HERV-H PROTFIN	Homo sablens butative brotein O-mannosultransferace (DOM/T2) DNA	Homo sepiens putetive protein C-mannosyltransferses (PON 12), IIIANA	Homo saplens DHHC1 protein (LOC51304), mRNA	Homo saplens MADS box transcription enhancer factor 2, polypeptide B (myocyte enhancer factor 2B) (MEF2B) mRNA	UI-HF-BN0-all-a-10-0-UI-r1 NIH MGC 50 Homo capiens CINA class IMA CE-2020803 El	UI HE-BNO-site - 10.01 II A NIM MOC EO Homo sentions control (MACE SOURCE)	601309466F1 NIH MGC 44 Homo sanlens CDN4 Alone IMA CE:3079867 5	AU130689 NT2RP3 Homo sapiens cDNA clone NT2RP2001282 F	TCAAP1E1219 Pediatric acute myelogenous leukamia celi (FAB M1) Baylor-HGSC project=TCAA Homo sapiens cDNA clone TCAAP1219	TCAAP1E1219 Pediatric acute myelogenous leukemia cell (FAB M1) Baylor-HGSC project≖TCAA Homo sablens cDNA clone TCAAP1219	Homo saplens chemokine MIP-2 gamma (MIP-2 gamma) mRNA complete cats	Homo saplens protein tyrosine phosphatase, non-recentor type 24 (PTDN)34 mpNA	Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA	Homo sapiens synaptojanin 1 (SYNJ1), mRNA	RC4-NT0057-160600-018-b05 NT0057 Homo saplens cDNA	CM3-UM0043-240300-127-e07 UM0043 Home saplens cDNA	CM3-UM0043-240300-127-e07 UM0043 Hamo saplens cDNA	CM3-UM0043-240300-127-e07 UM0043 Home septems aDNA	CM3-UM0043-240300-127-e07 UM0043 Homo saplens cDNA	or98e07.s1 NCL_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1603908 3'	ts89e07.x1 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE:2238468 3' similar to SW:PRO2_ACACAP19884 PROFILIN II:	Homo seplens placenta-specific 1 (PLAC1) mRNA	ym51h07.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:62071 5
ion i love oigi	Top Hit Database Source	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	N	TN	Į.	EST HUMAN	EST HUMAN	Т	Т		EST HUMAN	T				Γ	EST HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN		11496282 NT	П
5	Top Hit Acessian No.	AA297847.1	BE395061.1	8.0E-58 BE868715.1	58 AI798376.1	58 AI798376.1	11434921 NT	11434921 NT	7706132 NT	5174542 NT	58 AW 504109.1	7.0E-58 AW 504109.1	3E395061.1	6.0E-58 AU130689.1	38 BE242150.1	6.0E-58 BE242150.1		11434746 NT	11526291 NT	4507334 NT			П			5.0E-58 AA988183.1	1636745.1	5.0E-58 11496282	123072.1
	Most Similar (Top) Hit BLAST E Value	9.0E-58	9.0E-58	8.0E-58	8.0E-58	8.0E-58	8.0E-58	8.0E-58	8.0E-58	7.0E-58	7.0E-58	7.0E-58	6.0E-58	6.0E-58	6.0E-58	6.0E-58	6.0E-58	6.0E-58	6.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	6.0E-58 ₽	5.0E-58	5.0E-58	5.0E-58	5.0E-58 H23072.1
	Expression Signal	0.99	1.55	1.78	4.18	4.18	2.37	2.37	2.32	5.87	3.6	3.6	1.02	3.78	1.2	1.2	1.01	0.79	1.58	3.79	5.41	2.96	2.96	2.76	2.76	4.32	0.92	2:32	9.86
	ORF SEQ ID NO:	31297			26055		27301	27302			36773	36774	27718	27837	28316	28317	31824	36072		25740	26116	26589	26590	26589	26590	28727	29594		31829
	_ <u>w</u>	1		13355	13418	13418		14587	15740	23446	23529	23529	14979	15097	15668	15668	18857	22856	24754	13089	13469	13928	13926	13926	13926	16077	16970	18338	18863
	Probe SEQ ID NO:	5591	12516	575	639	639	1849	1849	2974	10762	10847	10847	2251	2375	2802	2902	6078	10208	12347	g	8	1122	1172	133	1173	3317	4229	5541	6085

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Page Page					Т	Т	\neg	т	Т		\neg	Т	Т	7	_	-т	_	Т	Ė	_	_	_	_	_	_	_	_			٠.					
Exan NO: ORF SEQ ID SIGNAL Expression Top Hit Acession Signal Most Similer Top Hit Acession Top Hit Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Top Hit Acession Top Hit Ace	oes Expressed in Brain	Top Hit Descriptor	Homo sabiens chromosome 21 segment HS21Chas	Homo sapiene enicel profein Yearnal a long III (ADV) - Data	Homo sapiens ribtin (NBS) mBNA complete out	Homo sapiens nibth (NBS) mBNA com-lete ada	Homo sapiens holocytochrome c synthese (cytochrome c homo homo) (ECCC) Exis	Homo sapiens hypothetical protein FI 1/0826 (FI 1/0828) mbNA	Homo sapiens mRNA for KIAA1617 protein nartial cde	Homo sablens ribonuclease 6 precursor (RNASFER) 1 mBNA	Homo saplens ribonuclease 6 precursor (RNASE6PI) mRNA	Homo saplens pre-mRNA splicing factor similar to S. cerevista Opera (DDD4s) DNA	Homo sapiens chromosome 21 segment HS21C018	Homo saplens mRNA for KIAA0641 protein martial cyts	Homo saplens mRNA for KIAA0611 protein partial cds	Homo saplens chimerin (chimeerin) 1 (CHN1) mRNA	Homo sapiens cat eve syndrome chromosome region candidate 4 (CECE4) BNA	Homo sapiens acetyl-Coenzyme A cerhovidese when ACACA with mining	Homo saplens Ran G Passe activating protein 1 (RANICAD4) monit	Homo saplens ATP synthase, H+ transporting, mitochondrial F1 complex O subjunit (Aliconomic).	conferring protein) (ATP50) mRNA	Homo saplens interleukin 10 receptor, beta (IL10RB), mRNA	-lomo saplens coagulation factor IX (plasma thromboplastic component, Christmas disease, hemophilla B) F9) mRNA	-luman beta-prime-adaptin (BAM22) gene expn 3	luman mRNA, Xq terminal portion	Homo seplens EGF-like repeats and discoidin Lilke domains 3 (FDI) 31 mRNA	Joino sapiens E18-55kDa-associated protein 5 / E18-AP51 mRNA	910e02.r1 Soares Infant brain 1NIB Home sapiens cDNA clone IMAGE:34862 8	lomo saplens peptide YY (PYY) mRNA	202185789F1 NIH MGC 45 Home sabiens cDNA clone IMAGE Annogra 5	02185789F1 NIH MGC 45 Homo saniens cDNA clone IMAGE 4300043 F	2V0-BT0702-170400-194-f09 BT0702 Homo sepiens cDNA	SC1TG081 normalized Infant brain cDNA Homo saniens cDNA clone c-14-08	V712977 DCA Homo sapiens cDNA clone DCAAZG04 5'	fomo saplens Serrunolevulinate symthase 2 (ALAS2) gene, complete cds
Exam ORF SEQ Expression Signal (Top) Hit Top Hit Acess No. Top Hit Acess No. 19074 32060 0.85 5.0E-58 AL163285.2 19074 32060 0.85 5.0E-58 AL163285.2 19074 32060 0.85 5.0E-58 AF051334.1 19697 32635 0.68 5.0E-58 AF051334.1 19697 32636 0.73 5.0E-58 AF051334.1 19698 32752 0.73 5.0E-58 AF051334.1 20945 33691 7.69 5.0E-58 AF051334.1 21918 35080 0.68 5.0E-58 AF04631.1 22408 35081 0.73 5.0E-58 AF04651.1 22408 37740 0.88 5.0E-58 AB01451.1 22647 36162 0.83 5.0E-58 AB01451.1 22647 36162 0.83 5.0E-58 A5046 25305 0.73 5.0E-58 A5046 25306 1.81 4.0E-58	gle Exon Pro	Top Hit Database Source	Į	LV.	Į.	L	LN LN	닐	N	N	N	NT.	N	 	TA L	7	5	Ę	5		-							Г				Γ	Г		П
Exon No: CNF SEQ Expression (Top) Signal ID NO: Signal Si	NIN NIN	Top Hit Acession No.		1330	AF051334.1	Ī	Š	8922693	Γ	5231227	5231227	11430647		Γ	Γ	1079	11526283	11426423	11418177		4502302	4504634	4503648			5031660 N	11424059 N		758981	Γ	Γ				
Exan ORF SEQ Express NO: NO: Signs NO: NO: Signs NO: NO: Signs NO: NO: NO: NO: NO: NO: NO: NO: NO: NO:		Most Similar (Top) Hit BLAST E Value	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58/	5.0E-58	5.0E-58	5.0E-58	5.0E-58	5.0E-58		4.0E-58	4.0E-58	4.0E-58	4.0E-58	4.0E-58	4.0E-58	4.0E-58	3.0E-58 R	3.0E-58	3.0E-58 B	3.0E-58 B	3.0E-58 B	3.0E-58 F	3.0E-58 A	20E-58 A
Exan SEQ ID NO: 19074 19074 19697 19698 220845 220847 220844 3 19689 2 18079 21808 23874 3 19898 23874 3 19898 23874 3 19898 2 2 19898 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 19898 2 2 198988 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898 2 2 19898		Expression Signal	0.85	1.61	0.68	0.68	0.73	7.69	0.7	0.68	0.68	0.88	1.78	0.83	0.83	2.69	1.81	1.5	2.67		4.5	0.88	1.09	1.7	1.03	1.25	7.06	2.67	2.36	2.78	2.78	0.63	1.43	1.4	11.9
				32147	32635	32636	32752	33691	34083	35089	32090	35614	35887	36161	36162	37740					25804	28212	26883	28091	28729	29113	37275		26791	28585	28586	31915	32114	32314	26350
Probe SEQ ID NO: NO: 6880 6880 6880 6880 7006 7006 7006 7006			19074	L.	Ĺ		l		1	1	' 1	ı	- 1		22947	24405	25305	25330	25001		13162	13551	14199	15349	16079	16476	23974	13127	14116	15937	15937	18944	192	19309	13686
		Probe SEQ ID NO:	8301	6379	9899	9899	7006	7869	8251	8239	9239	9757	100Z3	10300	10300	11819	12071	12512	12732		88	200	1452	2637	3319	3723	11315	326	1368	3174	3174	6167	6352	8 4	919

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		891 60S element binding				P:ZK328.1 SALCIUM									mRNA A															
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Be08b07.yf NIH_MGC_7 Homo sepiens cDNA clone IMAGE:2823733 5' similar to gb:X69391 60S RIBOSOMAL PROTEIN L6 (HUMAN); gb:X81987 M.musculus mRNA for TAX responsive element binding protein (MOLISE).	601499981F1 NIH MGC 70 Homo soutons above alera il MOT 30 Homo soutons ale	1601499961F1 NIH MGC 70 Homo sapiens CONA clore IMAGE 3601911 6	UI-H-BW1-ams-g-11-0-UI,s1 NCI CGAP Sub7 Homo sablens cDNA clone IMAGE-3071060 3	em57e02.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1838674 3' similar to WP:ZK328.1 GE05085 UBIQUITIN CONJUGATING ENZYMEY, RECOVERIN SUBFAMILY OF EF-HAND CALCIUM RINDING PROTEIN .	MO08h08 r1 Sogress feetal liver surface and School and	officers of the special interpretation of the second of th	Homo sapiens endondig recentor Endo/Bill Costo	Homo sablens endocytic receptor Endo180 (ENDO160) mKNA, complete cds	1601890812F1 NIH MGC 17 Himp soulest CENDO 160) mKNA, complete cds	hm25708 x1 NCI CGAP This Home seplent of the MACE secret of	Human complement component CS mBNA 2/2/3	Homo saniane NADIA dahudasasasa (i.i.l	EST369252 MAGE recent learners MAGD U	EST369252 MAGE resequences, MAGD Home septens curva	Homo saplens partial AF-4 dana Awas 2 to 7 and All months	Homo caplens stard requisitory element finding transcription forths of ACE TO TAKE	Homo sabiens G protein-counted recentor add (CPRsod) within	Homo saplens chondrolitin sulfate proteoglycen 2 (versionn) (CSPC3) mBMA	Homo sapiens chondroith suifate proteonlycen 2 (versicen) (CSDC2) mBN/A	oz43h01.x1 Soares NhHMPu S1 Homo sepiens cDNA clane IMACE:18784-20 3	RC1-BT0254-280100-015-601 BT0254 Homo sablens cDNA	Homo saplens hypothetical protein (LOC5126n) mRNA	EST385637 MAGE resequences. MAGM Homo sepiens cDNA	Homo sapiens myomesin (M-protein) 2 (165kD) (MYOM2) mRMA	AV751001 NPC Homo seplens cDNA clone NPCACH09 5'	209005.11 Sogres tests NHT Homo sablens CDNA close MAGE: 720,407 E	299105.r1 Sogres testis NHT Homo saplens CDNA clone IMACE 720407 g	Homo sapiens discs, large (Drosophila) homolog 2 (chapsyn-110) (DLG2), mRNA
igle Exon Pro	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	H IMAN	EST HUMAN	EST HUMAN	LN	LZ L	EST HUMAN	EST HUMAN	N	L L	EST HUMAN	EST HUMAN	LN	ŁN	L'A	ΙŻ	Ę	1	EST_HUMAN		EST_HUMAN		EST_HUMAN	EST HUMAN	T_HUMAN	
	Top Hit Acession No.	BE208532.1	BE907186.1	2.0E-58 BE907186.1		58 A1124874.1	58 R92567.1	2.0E-58 AI291407.1				2.0E-58 AW872641.1	Γ	6274549INT	58 AW957182.1	Τ		4759169 NT	5174444 NT	4758081 NT	4758081 NT		1.0E-58 BE061860.1	2031	1.0E-58 AW973537.1	4505314 NT			1.0E-58 AA412397.1 E	2994
	Most Similar (Top) Hit BLAST E Value	2.0E-58	2.0E-58	2.0E-58	2.0E-58	2.0E-581	2.0E-58	2.0E-58/	2.0E-58/	2.0E-58 /	2.0E-58	2.0E-58 /	1.0E-58 N	1.0E-58	1.0E-58 /	1.0E-58	1.0E-58	1.0E-58	1.0E-58	1.0E-58	1.0E-58	1.0E-58 A	1.0E-58 B	1.0E-58	1.0E-58 A	1.0E-58	1.0E-58 A	1.0E-58 A	1.0E-58 A	1.0E-58
	Expression Signal	10	3.4	3.4	1.12		8.0	1.12	2.83	2.83	21.77	2.43	0.86	2.41	1.81	1.61	1.13	2.37	1.6	0.88	0.88	4.75	1.31	9.0	9.0	99.0	0.91	0.66	0.68	121
	ORF SEQ ID NO:		30708			31771	31803	. 32511	32809	32810	36570	36813	26128	26464	26726	26727	26798	28251	27462	28938	28939	30256	31465	32633		34609	34714	34818	34819	35949
	Exon SEQ ID NO:	14016	Ιİ	25065	- 1	18811	18841	19489	19747	19747	23332	23565	13480	13805	14053	14053	14124	15510	14738	16282	16282	17841	18543	19508	20,708	21460	21571	21669	21669	22734
	SEQ ID NO:	1267	6273	5273	2966	6031	8062	6828	7058	7058	10841	10885	705	1046	1304	1304	1376	2805	2834	3628	3528	4913	6751	8764	8013	8768	88	8979	8979	100861

WO 01/57275

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The state of the s	Expression Signal (Top) Hit Acesslon Top Hit Acesslon Top Hit Descriptor Signal BLAST E Value Top Hit Descriptor Source	2.11 1.0E-58 X63392.1 NT H.sapiens immunoglobulin kappa light chain variable region L14	1.57	29.49	2.49	1.74 6.0E-59 BF035327.1 EST_HUMAN 601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	0.61 6.0E-59 AI750970.1 EST HUMAN cn06h02.y1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn06h02 random	1.32 5.0E-59 AW 157281.1 EST_HUMAN TR:O75786 O75786 GANGLIOSIDE-INDUCED DIFFERENTIATION ASSOCIATED PROTEIN 1.;	1.32 6.0E-59 AW157281.1 EST_HUMAN	6.98	6.55 5.0E-59 X83497.1 NT H.saplens DNA for ZNF8	u 1	4 04 E OF EO 44424720 NT	1.62 5.0E-59 AV762869.1 EST HUMAN	3.78 5.0E-59 11434908 NT	1.56	1.03	AF057720.1 NT	6.13 3.0E-59 AW 965524.1 EST HUMAN EST377682 MAGE resequences, MAGI Homo septens cDNA	7662247 NT		8.2	5.59 3.0E-59 AB029035.1 NT	5.59 3.0E-59 AB029035.1 NT	3.77 3.0E-59 4502014 NT	3.77 3.0E-59 4502014 NT Homo sepiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	1.45 3.0E-59	
	_=	L	L	Ĺ							L			L														20 20 0
	Exon SEQ ID ID NO:	24371	24404 37739	14953 27691	20774 33904	15536	20838 33970	14490 27189	14490 27190	15889 28530	17345 29978	47060				13548 26210	18249 31138	25238	12836	13030 25666						15891 28535	16557 29189	
	Probe SEO ID SE NO:	11780	11816	2225	8080	173	8144	1748 1	1748 1	3124 1		2000	ı	1	!	ı	5450 1	12203	9	219	1705 1	1705 1	ŀ	2125 1	į i	3126 1	3805 1	

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Single Exon Probes Expressed in Brain	Top Hit Database Top Hit Descriptor Source	27522 NT Homo sapiens protein thosphalasa recentor type T (BTDBT), mBNA				NT Human mRNA (or dbl proto-oncogene	NT Human mRNA for dbl prote-oncogene		NT H. sapiens CKII-alpha gene		EST HUMAN MR0-FT0144-260700-002-910 FT0144 Homo seplens cDNA	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN		Т	T_HUMAN	NT Homo sapiens alpha-tubulin mRNA, complete cds	EST_HUMAN 901176757F1 NIH_MGC_17 Hamo sapiens cDNA clone IMAGE:3631927 6'	ye25009.1 Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:118768 6' similar to SP:S21348 EST HUMAN S21348 HYPOTHETICAL PROTEIN 4 -		NT Homo sapiens Xdha mRNA for xanthine dehydrogenase, complete cds	CGAP_GCB1 Homo sepiens cDNA clone IMAGE:1309029 3' similar to TR:Q13537	NT	EST HUMAN	EST HUMAN			8849 NT	TN	EST_HUMAN
Single Exon Probes Ext	Top Hit Acession Database Source	7427522 NT Homo se	Z	3924074 NT	5464137 NT Homo sa	N	N	۲N	۲	1417868 NT	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	H H		EST_HUMAN	LN	EST_HUMAN	EST HUMAN	N	TN	FST HIMAN	NT	EST HUMAN	EST HUMAN	Т	11428849 NT Homo sap	8849 NT	TN	EST_HUMAN
	Most Similar (Top) Hit BLASTE Value	3.0E-59	3.0E-59 M95981.1	3.0E-59	3.0E-59	3.0E-59 X12556.1	3.0E-59 X12556.1	3.0E-59 X70251.1	3.0E-59 X70251.1	3.0E-59	2.0E-59 BF373329.1	2.0E-59 AA309774.1	2.0E-59 BF365554.1	2.0E-59 AW410698.1	2.0E-59 AW410698.1	2.0E-59 H61604 1		2.0E-59 AI631809.1	2.0E-59 L11645	1.0E-59 BE296411.1	1.0E-59 T92522	1.0E-59 D11456.2	1.0E-59 D11456	1 0F-59 AA748468 1	1.0E-69 AJ1308	1.0E-59 BE2568	1.0E-59 BE256814.1	1.0E-59	1.0E-59	1.0E-59	.1.0E-59 AJ130894.1	8.0E-60 AW977
	Expression Signal E	1.57	0.97	2.12	1.94	1.16	1.16	0.87	0.87	6.04	0.71	6.32	1.19	2.6	2.6	1.31		2.93	4.65	3.03	. 0.93		1.19	2.47		L		0.86	0.54	. 0.54	12.88	0.85
	ORF SEQ ID NO:	30143			33020	33653	33654	35794	35795				-	38663	36664	37274		31126	30719:		26964	27870	27871		33227	33392	33393	35210	35337	35338	33227	26178
	Exon SEQ ID NO:	17520	17713	ı		20527			- 1				23071	23421	23421	23970	1	24696	25263	12974	14276	15133	15133	15335	20135	20283	20283	22039	22157	22157	20135	13520
	Probe SEQ ID NO:	4789	.4890	6128	7259	7832	7832	9944	9944	12327	7693	9537	10425	10734	10734	11311	į	12091	12605	<u>8</u>	1529	2412	2412	2623	7462	7817	7617	9285	9504	9504	10760	747

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Single Exon Probes Expressed in Brain

	_	_	_	_		,	_		_	_			,		-	_	_						_					_			
Top Hit Descriptor	Homo sapiens small nuclear ribonucleoprotein D3 polypeptide (18kD) (SNRPD3) mRNA	Homo saplens differentiation-related gene 1 (nickel-specific induction protein) (RTP) mRNA	Homo sapiens differentiation-related gene 1 (nickel-specific induction protein) (RTP) mRNA	Homo sapiens mRNA for KIAA1081 protein, partial cds	hyaluronan-binding protein=hepatocyte growth factor activator homolog [human, plasma, mRNA, 2408 nt]	Homo sapiens phosphate cytldylytransferase 1, choline, beta isoform (PCYT1B), mRNA	Human mRNA for integrin alpha-2 subunit	Homo saplens S-antigen; retina and plineal gland (arrestin) (SAG), mRNA	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	Homo sapiens RAN binding protein 7 (RANBP7), mRNA	Homo sapiens chromosome 21 segment HS21C004	Homo sapiens chromosome 21 segment HS21C004	Homo sepiens MHC class 1 region	Homo sapiens MHC class 1 region	Homo saplens interleukin 10 receptor, beta (IL10RB), mRNA	Homo sapiens cullin 4A (CUL4A) mRNA, complete cds	Homo sapiens mRNA for KIAA0581 protein, partial ods	Homo sapiens omithine decarboxylase 1 (ODC1) mRNA	y1204.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:205087 6' similar to contains	vr12/04.r1 Spares fetal liver speen 1NFLS Homo sapiens cDNA clone IMAGE: 205087.5' similar to contains	LTR5 repetitive element;	601658751R1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886069 37	yq78h09.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:201963 5' similar to contains	OFR repetitive element;	wf52c07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2359212 3'	wf52c07.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2359212.3'	EST11498 Uterus Homo sapiens cDNA,5' end similar to similar to retrovirus-related pol	Institution NCI_CGAP_Kid11 Home sapiens cDNA clone IMAGE:3134913 3' similar to SW:RHOP_MOUSE October CTP.RHO RINDING PROTEIN 1	Hamo saplens chromosome 21 segment HS21C078	Homo sapiens v-raf-1 murine leukemia viral oncogene homolog 1 (RAF1), mRNA
Top Hit Detabase Source	L	FZ	TN	TN	ΕZ	F	Į.	ΤN	ΤN	NT	NT	NT	۱۶	N	Ę	N-	ΤN	LΝ	۲	NAME IN TOO	NIGHT ISS	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NAMI IL TOD	LN	NT
Top Hit Acession No.	4759159 NT	5174656 NT	5174656 NT	30 AB029004.1	S83182.1	-60 11420841 NT	S0 X17033.1	11428949 NT	11417118 NT	11417118 NT	5453997 NT	8.0E-60 AL163204.2	8.0E-60 AL163204.2	50 AF055068.1	7.0E-60 AF055066.1	4504634 NT	30 AF077188.1	30 AB011153.1	4505488 NT	7000011	1.100041.1	30 H58041.1	6.0E-60 BE964974.2				5.0E-60 AI807917.1	4.0E-60 AA289037.1	4 OF 80 BE106068 1	4L163278.2	4.0E-60 · 11433597 NT
Most Similar (Top) Hit BLAST E Value	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	8.0E-60	7.0E-60	7.0E-60/	7.0E-60	7.0E-60 /	7.0E-60 /	7.0E-60	100	7.05-00	7.0E-60	6.0E-60		6.0E-60	5.0E-60	5.0E-60	4.0E-60	1 05 80	4.0E-60	4.0E-60
Expression Signal	2.65	3.6	3.6	1.12	1.07	1.07	2.28	2.6	96.0	0.96	0.59	8.38	6.36	3.61	17.82	0.98	1.08	1.53	2.58	,	4.02	2.11	1.08		10.5	2.29	2.29	1.27	G C	0.88	1.29
ORF SEQ ID NO:		27632		31623	32178		33687	34675	35118		36348	36666	36667		26169	26228	27584		29527			37331	27639				25546		0.000	71000	37219
SEQ ID NO:			i '	18677		L		21529	21946	21946	23119	23423	23423	L	13511	13568	14855		16898			24027	14906	1		12908	12908	15738	10037	1	1 [
Probe SEQ (D NO:	1455	2169	2169	5892	6411	7598	7865	8837	9371	9371	10473	10736	10736	737	738	798	2124	2788	4158	1	3	11337	2177		8336	82	82	2972	7052	9024	11267

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	I was a market of the second o	From Septems War-I munifiered leukemia viral oncogene homolog 1 (RAF1), mRNA	001530440F1 NIF MGC_44 Homo sapiens cDNA clone IMAGE:3690395 5'	CONSCIPATION IN MINIMACE 44 HOMO Sepiens CDNA clone IMAGE:3890395 5'	Homo sapilens XII premidentification (FTD)	OVA-NN1140 25000 422 64 NN14425	RC3-LT0023-200400-423-101 INN149 Home sapiens cDNA	060h11.35 NOL CGAP_Kid3 Homo saplens cDNA clans IMAGE:1534053 5' similar to SW:UDP_MOUSE.	Lower Civilian In Contract (1)	Homo sapiens profine dehi-droppensos (regions of the control of t	ox56d09.x1 Sogres_NhHMPu_S1 Home sepiens cDNA clone IMAGE:1880337 3' similar to	SW:FORM_MOUSE Q05860 FORMIN;	Homo sapiens proline dehydrogenase (proline oxidase) (PRODH) mRNA	USE 1240227 FT NITE WISE BU HOMO Sapiens cDNA clone IMAGE:3830990 5'	Home sapiens Cel-102 protein (LOC57130), mRNA	abo7ho4.rf Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:840151 5' similar to contains	reconstruction of the second o	Hospiens 44,000 - 44,000 - 44,000 MRNA, complete cds; nuclear gene for mitochondrial product	Human bor protein mRNA. 5' end		Truito sapiens soutie carrier (SLC25A18) mRNA, complete cds; nuclear gene for mitochondrial product	Home series	Homo sapiens chromosome 21 infrasin mebila	nno1f12.y5 NCI_CGAP_Co9 Homo saplens CDNA clone IMAGE:1078485 5' similar to contains THR.tf THR	omo senieni,	Homo septions DNA polymerace mate catalife complete cds	Homo saplens corticotropin releasing homone recents 2 (ABLIBA). BMA	Homo saplens corticotropin releasing hormone receptor 2 (CRHR2) mRNA
gle Exon Pro	Top Hit Database Source	FN	FOT LINAM	EST LINAN	LA LA	LN	EST HIMAN	EST HUMAN	FOT HIMAN	L	L		HOMAN	HAMM	NUMBER		T HEJMAN	T			i.	LI BAAN	NICHOLL		E LIMANI	Т			
Sin	Top Hit Acession No.	11433597 NT	60 BE562611 1	BE562611 1	3.0E-60 6031190 NT	3.0E-60 AJ271735.1	3.0E-60 BF365143.1	3.0E-60 AW836196.1	30 A1792814 1	5174644 NT			3.0E-50 AIU40235.1 ES	į	120	11427120 NT			T		AY008285 1		7867	AF231919.1	A1794952 4		Γ	3044	4503044 NT
	Most Similar (Top) Hit BLAST E Value	4.0E-60	3.0E-60	3.0E-60	3.0E-60	3.0E-60	3.0E-80	3.0E-60	3.0E-60	3.0E-60	3.0E-60	. 10 6	3.05-50.	3.0F.80	3.0E-60	3.0E-60	3.0E-80	2 0F-80 4	2.0E-60 Z	2.0E-60 N	2.0F-60 A			2.0E-60 A	20E-60 A	2.0E-60 A	2.0E-60 A	2.0E-60	2.0€-60
	Expression Signal	1,29	4.44	4.44	1.92	1. 19.	0.57	2.12	-	4.97	4.97	28.0	4 32	0.47	1.26	1.26	2.08	383	7.35	1.28	- 65	1.98	0.69	0.73	0.86	1.87	96'0	2.43	2.43
	ORF SEQ ID NO:	37220		27306			30758	31260	30569	34132	34133	34310	34477	35398	37107	37108		25474	26831	27155	27166	28160	28969	29285	31962	32168	32393	30583	30584
	Exan SEQ ID NO:	23929	14590	l i	ı			18351	17933	20895	20995	21174	21333	22212	23829	23829	25297	12857	14151	14458	14467	15421	16321	16845	18983	19169	19378	17919	17919
	Probe SEQ ID NO:	11267	1852	1852	1862	4424	6294	5554	6856	8301	8301	8482	8841	9559	11162	11162	12686	8	1404	1715	1724	2714	3566	3895	6208	6400	9615	6750	6750

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Table 4
Single Exon Probes Expressed in

Noet Similar Signal Top Hit Acession Top Hit Top Hit Acession Signal Top Hit Acession Signal Top Hit Acession Top Hit Top Hit Acession Detabase Source Value T.06-61		
Noet Similar Top Hit Acession Top Hit	EST 14323 lests tumor Homo saplens cDNA 5' end EST 14323 Testis tumor Homo saplens cDNA 5' end Homo saplens hypothetical protein FLJ11028 (FLJ11028), mRNA	QV3-HT0513-060400-147-d01 HT0513 Home saplens cDNA QV3-HT0513-060400-147-d01 HT0513 Home saplens cDNA yv63d11.s1 Soares fetal liver spleen 1NFLS Home saplens cDNA clone IMAGE:248463 3' similar to
Expression Top) Hit Top Hit Acest Signal Top) Hit Top Hit Acest Signal Top) Hit Top Hit Acest No. 1,04 7,0E-61 7706 1,04 7,0E-61 7706 2,95 6,0E-61 BE408310.1 1,62 6,0E-61 BE408310.1 1,62 6,0E-61 BE408310.1 1,62 6,0E-61 BE257400.1 2,83 6,0E-61 AF19860.1 0,97 6,0E-61 AF19680.1 1,58 6,0E-61 AF19680.1 1,87 6,0E-61 AF096038.1 1,87 6,0E-61 AF096038.1 1,87 6,0E-61 AF096038.1 1,87 6,0E-61 AF096038.1 1,87 6,0E-61 AF0960388.1 1,135 6,0E-61 AF096038.1 1,135 6,0E-61 AF096038.1 2,22 5,0E-61 AF096038.1 1,73 5,0E-61 AF096038.1 2,56 5,0E-61 AF096038.1 3,38 5,0E-61 AF0778 <	T HUMAN	EST_HUMAN
Nosession (Tracess	2823	
Sygnes Significant Control of Con	3.0E-61	2.0E-61 BE16841
	0.64	0.82
	34428 25906 26607	26608
_ 	13271	13942 13942
Probe SEQ ID NO: 783 1288 1674 11474 11474 11474 11474 11474 11474 11474 11474 1168 116	8596 486 1190	1190

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חוקום באספס באלים הספס באלים הספס ביויסים ביוסים ביוסים ביוסים ביוסים ביוסים ביוסים ביוסים ביוסים ביוס	Top Hit Descriptor	yy03f11.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:270189 5'	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) non-catalytic accessory protein 1A (110/116kD) (ATP6N1A), mRNA	AV694317 GKC Homo saplens cDNA clone GKCELG06 5	Homo sapiens mRNA for KIAA0536 protein, partial cds	UI-HF-BN0-akd-f-12-0-UI.r/ NIH_MGC_50 Homo sapiens cDNA done IMAGE:3076774 5'	Homo sapiens polymerase (RNA) III (DNA directed) (38kD) (RPC39), mRNA	Homo sapiens ribosomal protein L44 (RPL44), mRNA	Homo sapiens origin recognition complex, subunit 2 (yeast homolog)-like (ORC2L) mRNA	Homo sapiens zona pellucida glycoprotein 3A (sperm receptor) (ZP3A), mRNA	xn11b09.y1 NCI_CGAP_LI5 Homo sapiens cDNA clone IMAGE:2693369 5' similar to contains element MSR1 repetitive element.	601273513F1 NIH_MGC_20 Hamo sapiens cDNA clone IMAGE:3614687 5'	Homo sapiens KIAA0806 gene product (KIAA0806), mRNA	QV2-HT0577-140300-077-g06 HT0577 Homo sapiens cDNA	Homo sapiens TRAF family member-associated NFKB activator (TANK) mRNA	Homo sapiens TRAF family member-associated NFKB activator (TANK) mRNA	UI-H-BW0-alt-b-08-0-UI.s1 NCI_CGAP_Sub6 Hamo sapiens cDNA clone IMAGE:2732871 3'	UI-H-BW0-qlt-b-08-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2732871 3'	Hamo sapiens chromosame 21 segment HS21C010	H.sapiens carbonic anhydrase VII (CA VII) gene, exons 4,5,6, and 7, and complete cds	Homo sapiens KIAA0783 gene product (KIAA0783), mRNA	Homo sapiens survival of motor neuron 1, telomeric (SMN1), mRNA	Human P40 T-call and mast call growth factor (hP40) gene, complete cds	Homo saplens SC35-Inferacting protein 1 (SRRP129), mRNA	Homo sapiens hypothetical protein FLJ20128 (FLJ20128), mRNA	Homo sapiens hypothetical protein FLJ20128 (FLJ20128), mRNA	Homo sapiens growth hormone releasing hormone (GHRH), mRNA	Homo sapiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3	(UBE2D3) genes, complete cds	MR0-BN0070-040400-010-h01 BN0070 Homo sapiens cDNA	Homo sapiens cadherin 18 (CDH18), mRNA	Homo sapiens KIAA0971 protein (KIAA0971), mRNA	Homo sapiens actinin, alpha 4 (ACTN4), mRNA
מסו וווסעד סול	Top Hit Database Source	EST_HUMAN)		-HUMAN		L HUMAN					EST HUMAN	Г		EST_HUMAN (T_HUMAN												T_HUMAN			
5	Top Hit Acession No.	-61 N39397.1	11426168 NT	-61 AV694317.1	-61 AB011108.1	-61 AW 500256.1	11421778 NT	11419729 NT	5453829 NT	E8005083 NT	61 AW827281.1		62319	61 BE174455.1	4759249 NT	4769249 NT	-61 AW298181.1		-61 AL163210.2		7662303 NT	1416891	1.0E-81 M30135.1	4759171 NT	8923130 NT	8923130 NT	11034840 NT			-61 AW999726.1	11416280 NT	11428892 NT	11425578 NT
	Most Similar (Top) Hit BLAST E Value	2.0E-61	20E-81	2.0E-61	20E-61	2.0E-61	2.0E-61	2.0E-61	1.0E-61	1.0E-61	1.0E-61		1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-81	1.0E-61	1.0E-61	1.0E-91	1.0E-61		1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61
	Expression Signal	1.04	0.98	0.98	0.99	1.67	2.3	1.81	1.11	3.71	1.42	1.47	0.86	1.2	0.81	0.81	8.11	8.11	0.75	1.82	0.79	1.29	7.11	0.67	1.42	1.42	3		3.06	2.7	7.73	5.24	2.84
İ	ORF SEQ ID NO:		32090			35676	36014		26188	27304	27858	28257	28786	29106	29773	29774	30157	30158	30247	30772	31310	31511	32482	32732	32845	32846	33861		34041		35110		H
	SEQ ID NO:	15357	19102	21604	i .		22798	23482	13528	14589	14922	15607	16128	16468	17144	17144		17535	17632	18114	H		19461	19684	19780	19780	20728			21852	21936	1	
	Probe SEQ ID NO:	2647	6332	8913	9462	9822	10150	10799	756	1851	2183	2839	3369	3715	4407	4407	4804	4804	4905	5309	5603	5783	0890	6991	7091	7091	8033		8212	9182	9257	8558	10531

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Single Exon Probes Expressed in Brain	Top Hit Acession Top Hit Descriptor Source	FIN		AGO NT			=084386.1 EST HUMAN IRC4-BT0310-110300-045-40 BT0340-H	FOU	EST HIMAN	The state of the s	140708E NIT	N O O O O O O O O O	EST HOMAN	ODEE NIT	EQT LIBRAL	EST HIMAN	EST HIMAN	39 NT	6.0E-62 AW814393.1 EST HUMAN MR3-ST0203-130100-025-00 ST0303 Home	EST HUMAN	N-	LN L	Z	LN.	1758 NT	TOT LAND	6758 NT	FST HIMAN	574	
}		1 ABOA45	1 AROUZE	9			BE0643	62 44830420 4	62 AV714334.1	D17480	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	TOE BO A LOSSON	1004101		6.0E-62 A17R2R01 1	6.0E-62 AI762801 1	AW50112		AW81436	A1950528.	AJ271735	AJ271735	U39487.1	U39487.1	5.0E-62 4506	AA431093 1	4	AW410687 1	٤	E
	Most Similar (Top) Hit BLAST E Value	1.0E-6	101-8	1.0E-6	1.0E-6	1.0E-61	9.0E-6	8.08	7.0E-62	7 OF 65	7.0E-82	1 0 1	8 OF -83	8 0 1 2	8.0E-82	6.0E-62	6.0E-62	6.0E-62	6.0E-62	5.0E-62	5.0E-62	6.0E-62	5.0E-62	5.0E-62	5.0E-62	5.0E-62				5.0E-62
	Expression Signal	1.40	1.83	3.02	3.02	11.58	1.45	1.1	1.62	0.74	0.64	7	1.42	4	3.03	3.03	0.72	1.45	3.27	2.8	4.25	4.26	1.35	1.35	2.92	2.6	0.55	6.45	2.85	2.85
	ORF SEQ ID NO:	36780			30727	30988	36113	29885	26501	28907	31547	37347			33289	33290		33981	36108	25840	27863	27864	28048	28049	28820	29680	34278	35267	37180	37181
	Exon SEQ ID NO:	23535	L			. 1	22903	17249	13843	16263	18615	24014	15764	16138	20196	20196	20679	20849	21933	13192	16127	15127	16312	15312	16171	17032	21139	22095	23894	23894
	SEQ ID NO:	10855	11006	12007	12007	12659	10266	4514	1085	3497	5826	11323	2998	3379	7625	7525	7984	8155	9254	407	2408	2400	2698	2598	3413	4293	8447	9417	11231	11231

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						21 - 1 1 1 1 2 1 1 1 2 1 1 1 1 1 1 1 1 1	ישור באכון ו הפסי באף הסיים וו הישווי
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
820	13591	26258	1.95	4.0E-62	AW161479.1	EST_HUMAN	au71d03.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);
820	13591	26259	1.95	4.0E-62	AW 161479.1	EST_HUMAN	au71d03.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);
821	13591	26258	2.96	4.0E-62	AW 161479.1	EST_HUMAN	au71403.y1 Schnelder fetal brain 00004 Homo saplens cDNA clone IMAGE:2781701 5' similar to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);
821	13591	26259	2.98	4.0E-62	AW161479.1	EST_HUMAN	au71d03.y1 Schneider fetal brain 00004 Homo saplens cDNA clone IMAGE:2781701 5' similær to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);
2459	15177	27916	1.78	4.0E-62	AI827900.1	EST_HUMAN	W12b08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2350359 3' similar to gb:X57138_ma1 HISTONE H28.2 (HUMAN);
2459	15177	27917	1.78	4.0E-62	AI827900.1	EST_HUMAN	W12b08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2350359 3' similar to gb:X57138_ma1 HISTONE H2B.2 (HUMAN);
3394	16153		6.34	4.0E-62	LN 288294	IN	Homo saplens keratin 18 (KRT18) mRNA
5833	18622	31555	1.84	4.0E-62	4506978 NT	NT	Homo sapiens solute carrier family 13 (sodium-dependent dicarboxylate transporter), member 2 (SLC13A2) mRNA
6204	18979	31958	1.9		11420654 NT	IN	Homo sapiens ubiquitin specific protease 9, X chromosome (Drosophila fat facets related) (USP9X), mRNA
7071	19782	32826	1.84	4.0E-62	11421041 NT	۲۸	Homo saplens phosphoribosyl pyrophosphate synthetase 2 (PRPS2), mRNA
7534	20204		2.48	4.0E-62	7657057 NT	ΗN	Homo sapiens eukaryotic translation initiation factor 2B, subunit 2 (beta, 39kD) (EIF2B2), mRNA
7534	20204		2.48	4.0E-62	7657057 NT	TN	Homo saplens eukaryotic translation initiation factor 2B, subunit 2 (beta, 39kD) (EIF2B2), mRNA
8071	20765		1.06	4.0E-62	11429973 NT	ΝΤ	Homo sapiens 26S proteasome-associated pad1 homolog (POH1), mRNA
8745	21437		4.97		AB033089.1	Į.	Homo sapiens mRNA for KIAA1263 protein, partial cds
10934	23614	_		4.0E-62	Z78766.1	N	H.sapiens flow-sorted chromosome 8 HindIII fragment, SC8pA16D3
10934	23614			4.0E-62	Z78766.1	占	H.sapiens flow-sorted chromosome 6 Hindlil fragment, SC6pA16D3
11995	24533		2.81	4.0E-62		·	Homo sapiens putative nuclear protein (HRIHFB2122), mRNA
12590	24947		1.34	4.0E-62		M	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA
12645	24942		16.72	4.0E-62	11417862 NT	NT	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA
12645	24942	30982	16.72	4.0E-62	11417862 NT	NT	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA
12692	24976		2.72	4.0E-62	11430460 NT	TN	Homo saplens low density lipoprotein-related protein 2 (LRP2), mRNA
72	12899	25535	0.89	3.0E-62	4557794 NT	TN	Homo sepiens neurofibromin 2 (bilateral acoustic neuroma) (NF2) mRNA
3041	15807	28452	1.11	3.0E-62	AB040909.1	TN	Homo sapiens mRNA for KIAA1476 protein, partial cds
3041	15807		1.11	3.0E-62	9.1	TN	Homo saplens mRNA for KIAA1476 protein, partial cds
3686	16439	29081	5.41	3.0E-62	X52858.1	N	Human cyclophillin-related processed pseudogene

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Table 4
Single Exon Probes Expressed in Brain

				_	_			_	_			_	_																		
URIA III DESCRITI	Top Hit Descriptor	wa33104.x1 NCL_CGAP_Kid11 Homo sapiens oDNA clone IMAGE:2299903 3' similar to contains THR t2	Ink repetitive element;	From Sapiens chromosome 21 segment HS21C084	RC0-BN0284-300500-031-e05 BN0284 Homo sapiens cDNA	Homo sapiens mannosidase, beta A, Iyeosomai (MANBA) gene, and ubiquitin-conjugating enzuma E2D 3	(UDE ZU3) genes, complete cds	4.44-51.025/-081199-017-e03 BT0257 Homo sapiens cDNA	Holling septems intersectin 2 (SH3D1B) mRNA, complete cds Holling semions ADD/ATB	effort.1.1 Soares NhHMPu_S1 Home saplens cDNA clone IMAGE:1047404 5' similar to WP:K01H12.1	1/201508104 14 Edg (mmonth 19 John 19	OVO LTAKES SEGMENT SET 1 WIND THE SECOND CONE OK FZP568F104 6	Home control to the c	History Septens hypothesical protein PLJ20212 (FLJ20212), mRNA	Trainian glucagon-like peptide-1 receptor (GLP-1) mRNA, complete cds	protein LYBB (RPL189), Ca2+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR)	COM protein (CDM), adrenoleukodystrophy protein >	abooco.s. stratagene reta retina 937202 Homo sapiens cDNA clone IMAGE:839906 31	2803110:S1 Societe Tetal_neart_NbHH19W Homo sepiens cDNA clone IMAGE:409771 3	280910.31 30ales letal near NbHH19W Home saplens cDNA clone IMAGE:4097713'	Homo saplens KIA ANTRA gene product (VIA A ARTES)	Homo saplens KIAA0789 gene product (N.) A 0700.	Sablens Nacional acid about the	Hearless by Secure and prospirates gene (EC 3.13.2) Exon 9	and Address to the Control of the Co	H. sanlans from social of the contro	Proceedings and cilionosome 6 Hindill fragment, SC6pA14D8	Homo captons control of the Component Krp46 (LOC56915), mRNA	Homo septents cannerin Edir LAG seven-pass G-type receptor 1 (CELSR1), mRNA	OVA ST024 181400 027 4/15 ST0221 1	C18159 Human placents cDNA (TFilingers) Home sapiens cDNA
מון וומעדורו	Top Hit Database Source	TOD TOD	NA TOWN	EST HIMANI		ļ.	1444	Т	LZ LZ	H	T	Т	- CANO			<u> </u>	H. IMANI	Т	Т	Т	Т				T HI MAN					T HIMAN	HUMAN
	Top Hit Acession No.	-62 AIR32733 1		Τ		82 AE224een 4	1	T	T	2			320	82 23503.1		32 U52111.2	-		Τ	T	32289	7662289 NT	15533.1 NT				1424055	11418322 NT	11430460 NT	AW816405.1	
	Most Similar (Top) Hit BLAST E Value	3.0F-82	2.0E-82	2.0E-62	2.0E-82	205.82	2 OF-82	1.0E-82	1.0E-62	1.0E-82 /	1.0E-62/	1.0E-82	1.0E-62	1.0E-62		1.0E-82[C	1.0E-62 A	1.0E-62 A	1.0E-62 A	1.0E-62 A	1.0E-62	1.0E-62	1.0E-62 X15533.1	1.0E-62 X15533.1	1.0E-62 AA465170.1	1.0E-62 Z78698.1	1.0E-62	1.0E-62	1.0E-62		9.0E-63 C
	Expression Signal	5.82	2.36	4.89	4.89	3.8	4.81	1.87	11.01	1.04	66.0	0.71	1.57	6.0		0.86	0.91	2.94	2.94	0.5	2.13	2.13	2.02	202	3.54	2.01	1.52	2.25	2.99	2.59	1.53
	ORF SEQ ID NO:	34267			34512			26443	26970	27239	28328		29855	30405		31948	32782	32796	32797	34491	34797	34798	34838	34839	35177	37333	37792		30890	25765	
	Exon SEQ ID NO:	21130	13960	21365	21365	22720	24284	13781	14283	_	ľ	- 1	17226	17790		18972	19726	19736	19736	21347	21647	21647	21689	21689	22007	24029	24460	24838	24966	13130	15068
L	Probe SEQ ID NO:	8438	1209	8673	8873	10072	11689	1021	1536	1791	2915	4317	4490	6071		6196	7034	7045	7045	8655	8928	8856	6668	688	9457	11339	11866	12474	12673	329	2345

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Table 4
Single Exon Probes Expressed in Brain

Single Later Lobes Lapressed in train	xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx	29395 7.42 9.0E-63 AB002348.2 NT	29396	37798 3.05	30869 1.63 9.0E-63 Y15056.1 NT	32837 3.86 9.0E-63 11426985 NT	33501 0.91 9.0E-63 4885544 NT	34055 1.38	36736 2.03 9.0E-63 7662289[NT	36737 2.03	27803 1.32 8.0E-63 4557734 NT	27834	28861 3.02 8.0E-63 AF198349.1 NT	28862	29600 3.31 8.0E-63 AL163268.2 NT	2.09	48.05 6.0E-63 AA420803.1	34612 1.97 5.0E-63 11526464	28726 0.84 4.0E-63 AL1632	29174 1.16 4.0E-63 AB014607.1 NT	29175 1.16 4.0E-63 AB014607.1 NT	32115 3.64 4.0E-63 AW 750372.1 EST_HUMAN	32116 3.84 4.0E-63 AW 750372.1 . EST_HUMAN	37021 2.3 4.0E-63 AW134709.1 [EST_HUMAN	37022 2.3 4.0E-63 AW134709.1 EST_HUMAN	37771 4.32 4.0E-63 AA362834.1 EST_HUMAN	27377 2.82 3.0E-63 AB018260.1 NT	28225 2.26 3.0E-63 J00310.1 NT	26636 11.81 3.0E-63 6005963 NT	32150 32.78 3.0E-63 11545810 NT	35444 1.15 3.0E-63 BE876158.1 EST_HUMAN	35446 1.16 3.0E-63 BE876158.1 EST_HUMAN	25639 1.09 2.0E-63[U07804.1
																								L				L					
	D SEO ID NO:	4020 16766	16766	5164 17895		7082 19772		224 20918	10816 23499	1	15066	l	154 16210		4234 16975	l	l	ı	3315 16075	788 16540	<u> </u>				777 23747	L	_	L	2824 13967	6382 19151		9605 22258	
	Probe SEQ ID NO:	4	4	Ċ	, R	_		Ö	Ę	ê	, i	K	6	્ર	<u> </u>	Ľ	58	ω	<u></u>	్	<u>س</u>	ශ්	త	11077	Ĕ	Ē	Ĕ	Ñ	T _K	8	ග්	ð	Ĺ

Page 334 of 536 Table 4 Single Exon Probes Expressed in

		T	7_	Т	T	T	Т	T	_,	_	T	_	т	_				 -	_	_	_		-		_	-	-, -
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens eyes absent (Droscophila) homolog 2 (FYA2) mRNA	Homo sapiens glutamate-cysteine ligase (gamma-glutamycystaine synthetase), catalytic (72.8kD) (GLCLC) mRNA	Homo sapiens Down syndrome candidate region 4 (DSCB4) — Byla	Homo sapiens RHCE mRNA for Rh blood CE grain softwar actions the	Homo saplens RHCE mRNA for Rh blood CE group entire maintenantal	601301627F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638403 F1	Homo sapiens amyori beta (A4) previnces prodein (protection)	Homo sanians chromosome 3 sushalomoda main (processe nextra), Alznemer disease) (APP), mRNA	Homo sapiens polycystic kidney disease-associated protein (DKD1) rene commissions.	Homo saniare im dimediados mestis.	Home saplens dutaming hearing engineering gene, partial cas, cros gene, complete cas, and unknown gene	Homo septens similar to ectonucleotide pyrophosphatase/phosphodiesterase 3 (H. saplens) (LOC63214), MRNA	0V1-ET0170-040700-28E-08 ET0470 U	OVI-FT0170-040700-098-508 FT0170 II	Homo saniens providing Values of MB 40000 100000 100000 100000 1000000	Homo saplens protein kinase, cAMP-dependent, regulatory, type II, beta (PRKAR2B), mRNA	Human germiline T-cell receptor beta chain Dopamine-beta-hydroxylase-like, TRY1, TRY2, TRY3, TCRBV27S1P, TCRBV22S1AZN1T, TCRBV6S1A1T, TCRBV6S1A1T, TCRBVSS1A1T, TCRBV13S3, TCRBVSS7P, TCRBV7S3A2T, TCRBV13S3, TCRBV6S7P, TCRBV7S2A1N4T	TCRBV13S9/13S>	TOTAL ORDINAL MICH. DATA	Troillo sapiens mild. Mr.N.A., partial cds	Homo saniens Carbonic annydrase-related protein 10 (LOC56934), mRNA	Homo sapiens mRNA for KIAA1624 profein partial cds	Homo espiens similar to sema domain, Immunoglobulin domain (tg), short basic domain, secreted, (semenonin) 3A (H. sapiens) (1.0003333) PRNA	Homo sapiens chromosome 21 segment HS21C310	Homo saplens kinesin family member 3B (KIF3B), mRNA	Homo sapiens kinesin family member 3B (KIF3B), mRNA
igle Exon Pro	Top Hit Database Source	NT	LN	۲	N-I	N	EST_HUMAN	IN	Į.	ĮN	F	LN		T HI IMAN	Т				L L						±N		
נֿה 	Top Hit Acession No.	4885226 NT	4557624 NT	7657042 NT	AB030388.1	AB030388.1	2.0E-63 BE410739.1	4502166 NT	-63 AF109718.1	-63 L39891.1	2.0E-63 AF111167.2	2617	11419429 NT	2.0E-63 BF373541.1	Γ	2.0E-63 11421940 NT	11421940 NT		-63 U68059.1	T	0388	9910365 NT	-63 AB046844.1	11421514 NT	63 AL163210.2	3949	11420949 NT
	Most Similar (Top) Hit BLAST E Value	2.0E-63	2.0E-63	2.0E-63	2.0E-63	2.0E-83	2.0E-63	2.0E-63	2.0E-83	2.0E-83	2.0E-63	2.0E-63	2.0E-63	2.0E-63	2.0E-63	2.0E-63	2.0E-83	!	2.0E-63 L	2 0F-83 A	2 0F-83	2.0E-83	2.0E-63 A	2.0E-63	2.0E-63 A	2.0E-63	2.0E-63
	Expression Signal	1.68	2.34	5.57	1.43	1.43	1.1	3.44	2.02	3.74	-	96.0	1.26	2.96	2.96	0.84	0.84		1.67	0.88	1.45	1.45	0.87	0.56	3.96	1.35	1.35
	ORF SEQ ID NO:	25647		26244	26994	26995	27203	28563	28690	29274	30169	30435	30505	31512	31513	31837	31838		32425	32426	32477	32478	33454	33490	34262	34791	34782
	Exon SEQ ID · NO:	13006	13270			_	14502			16835	17644	17818	25062	18585	18585	18871	18871	1000	19411	19411	19468	19456	20342	20377	21124	21643	21843
	Probe SEQ ID NO:	193	485	807	1559	1659	. 1760	3154	3279	3885	4813	6609	5181	6794	6794	6093	S609	680	6849	6849	6975	9769	7678	7713	8431	8952	8962

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					6		
Probe SEO ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9839	22490	35691	1.12	2.0E-63	AL 163218.2	TN	Homo sapiens chromosome 21 segment HS21C018
10647		36577	6.43	2.0E-83	N78945.1	EST_HUMAN	zb18b05.s1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA cione IMAGE:302385 3' similar to gb:X17208 40S RIBOSOMAL PROTEIN S4 (HUMAN);
10672	_	38604	2.96	2.0E-83	AF099810.1	LN	Homo sapiens neurexin III-alpha gene, partial cds
10672	23363	38605	2.98	2.0E-63	AF099810.1	NT	Homo sapiens neurexin III-alpha gene, partial cds
12098	25177	30807	5.97			1	Homo sapiens aconitase 2, mitochondrial (ACO2), mRNA
1502	14248	L	1.28		F08485.1	EST_HUMAN	HSCZVD/11 normalized Infant brain cDNA Homo sapiens cDNA clone c-zvd11
1502	ı	26935	1.28	1.0E-63	F08485.1	EST_HUMAN	HSCZVD111 normalized infant brain cDNA Homo sapiens cDNA done c-zvd11
4308	17047	29672	2.92	1.0E-83	F08485.1	EST_HUMAN	HSCZVD111 normalized Infant brain cDNA Homo sapiens cDNA clone c-zvd11
4308	17047		2.92		F08485.1	EST HUMAN	HSCZVD111 normalized infant brain cDNA Homo saplens cDNA clone o-zvd11
5268	18074		8.0	1.0E-63	AJ271736.1	NT	Homo sapiens Xq pseudoautosomal region; segment 2/2
5683	18476	31394	1.4		AW 582266.1	EST_HUMAN	QV0-ST0215-060100-083-b09 ST0215 Home sapiens cDNA
6298	19071	L	0.68		AW451950.1	EST_HUMAN	UI-H-BI3-alt-h-02-0-UI.s1 NCI_CGAP_Sub5 Horno sapiens cDNA clone IMAGE:3068763 3'
6298	19071	32056	99.0	1.0E-63	AW451950.1	EST_HUMAN	UI-H-BI3-alt-h-02-0-UI.s1 NCI_CCAP_Sub5 Homo sapiens cDNA clone IMAGE:3068763 3'
8371	21084		2.68	1.0E-63	AL163247.2	NT	Homo saplens chromosome 21 segment HS21C047
12737	25286		4.04	1.0E-63	AL163207.2	NT	Homo sapiens chromosome 21 segment HS21C007
7766	20462	33586	4.36		AI478186.1	EST_HUMAN	tm50b07.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2161525 3
1024	13784		6.16		BE280796.1		601155232F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3139038 5
6049	18829	31792	3.88		BE885755.1	EST_HUMAN	601508968F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3910338 5
11916	24479		7.34		11418177 NT	LN	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA
11970	24516		1.6		T60651.1	EST_HUMAN	yb98b02.r1 Stratagene lung (#937210) Homo sapians cDNA clone IMAGE:79179 5
3520	16276		1.13		BE39432	EST_HUMAN	601311455F1 NIH_MGC_44 Homo sepiens cDNA clone IMAGE:3633204 5'
4683	17417		2.73			님	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA
4683	17417	30053	2.73		4507490 NT	F	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA
9933	22581	35779	3.43	7.0E-64	Y07848.1	۲	Homo sapiens EWS, gar22, rrp22 and barn22 genes
1718	14459	27156	1.63	6.0E-64	Al851992.1	EST_HUMAN	wb51e07.x1 NCI_CGAP_GC6 Homo saplens cDNA clone IMAGE:2309220 3' stmilar to gb:M15182 BETA- GLUCURONIDASE PRECURSOR (HUMAN);
1718	1				_	EST HUMAN	wb51e07.x1 NO_CGAP_GC8 Homo septens cDNA clone IMAGE:2309220 3' simitar to gb:M15182 BETA- GLUCURONIDASE PRECURSOR (HUMAN);
3120	ı	L				EST_HUMAN	wv13e03.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2529436 3'
3120			4.39		AW026445.1	EST_HUMAN	wv13e03.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2529436 3'
5534	18332	31237	2.46			K	Homo sapiens MCP-1 gene and enhancer region
5534	18332			6.0E-64	Y18933.1	닐	Homo sapiens MCP-1 gene and enhancer region
5555	18352	31261	4.08		M13975.1	¥	Homo saplens protein kinase C beta-il type (PRKCB1) mRNA, complete cds

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Table 4

Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo saplens atrophin-1 interacting protein 1; activin receptor interacting protein 1 (KIAA0705), mRNA	Homo sapiens calcitonin receptor (CALCR), mRNA	Homo saplens calcitonin receptor (CALCR), mRNA	Homo sapiens mesenchyme homeo box 1 (MEOX1), mRNA	Homo sapiens mesenchyme homeo box 1 (MEOX1), mRNA	Homo sapiens acetyl-CoA synthetase (LOC55902), mRNA	Homo sapiens progressive ankylosis-like protein (ANK) mRNA, complete cds	trkC [human, brain, mRNA, 2715 nt]	Homo sapiens stromal antigen 3 (STAG3), mRNA	Homo sapiens stromal antigen 3 (STAG3), mRNA	wv13e03.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2529438 3'	wv/3e03.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2529438 3'	Homo sapiens Interleukin 10 receptor, beta (IL10RB), mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo sapiens phosphoglucomutase-related protein (PGMRP) gene, complete cds	Homo sapiens phosphoglucomutase-related protein (PGMRP) gene, complete cds	Human (3)mbt protein homolog mRNA, complete cds	Homo sapiens KIAA0618 gene product (KIAA0618), mRNA	Homo saplens KIAA0618 gene product (KIAA0618), mRNA	Homo saplens putative transcription factor CR53 (CR63) mRNA, partial cds	Horno sapiens mRNA for KIAA0903 protein, partial cds	601590382F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3944397 5'	RC3-ST0197-120200-015-a03 ST0197 Homo saplens cDNA	RC3-ST0197-120200-015-a03 ST0197 Homo sapiens cDNA	C18895 Human placenta cDNA (TFujiwara) Homo saplens cDNA clone GEN-569E02 5'	601589565F1 NIH_MGC_7 Home sapiens cDNA clone IMAGE:3843577 5'	AV711714 DCA Homo sapiens cDNA clone DCAAMC01 5'	AV711714 DCA Homo sepiens cDNA clone DCAAMC01 5	H.sapiens isoform 1 gene for L-type calcium channel, exon 28	RC6-FN0019-280600-011-G11 FN0019 Homo sapiens cDNA	Homo sapiens golgi matrix protein GM130 (GOLGA2) mRNA, complete cds
le Exon Probe	Top Hit Database Source							TN.	FZ	±N			EST_HUMAN W					FA	TN T	F						П			EST_HUMAN 6		T. HUMAN	П	T_HUMAN	T I
Sing	Top Hit Acession No.	6912461 NT	11422189 NT	11422189 NT	11525879 NT	11525879 NT	11420555 NT	6.0E-64 AF274753.1	S78475.1	11420197	11420197 NT	ŀ	6.0E-64 AW026445.1	11526198 NT							7662205 NT	32205					3.1		3.0E-64 BE794381.1				3.0E-84 BF370000.1	7
	Most Similar (Top) Hit BLAST E Value	6.0E-64	6.0E-64	6.0E-64	6.0E-84	8.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	6.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-64 L40933.1	5.0E-64 L40933.1	5.0E-64 U89358.1	5.0E-64	5.0E-64	5.0E-64 /	5.0E-64	4.0E-64	4.0E-64	4.0E-64	3.0E-64 C18895.1	3.0E-64 E	3.0E-64	3.0E-64	3.0E-64 Z26273.1	3.0E-84	3.0E-84 p
	Expression Signal	1.26	0.62	0.62	2.34	2.34	9.76	2.09	2.78	6.01	6.01	1.84	1.84	4.97	2.85	2.85	1.84	1.3	1.3	1.37	4.85	4.85	6.71	1.05	0.58	2.23	2.23	5.41	0.89	2.22	2.22	1.21	3.34	1.93
ŀ	ORF SEQ ID NO:	31269	31453	31454	32889	32890	35077	35240	35458	36600	36601	28524	28525	31089	26235	26236	26738	26827	26828	27149	26897	26898	29328	29455	33493	36644	36645	27658	28662	28841	28842	31734	32169	34198
	Exon SEQ ID NO:	18361	18531		L	19823	21905			23360			15885	24608	13573	13573	14064	14148	14148	14449	14210	14210	16690	16828	20380	23404	23404	14924	16011	16192	16192	18771	19170	21058
	Probe SEQ ID NO:	5564	6739	5739	7136	7136	9226	9406	9618	10669	10669	10941	10941	12115	801	801	1316	1401	1401	1706	2829	2829	3940	4085	7716	10715	10715	2195	3249	3436	3436	2990	6401	8365

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Table 4
Single Exon Probes Expressed in Brain

WO 01/57275

Single Exon Probes Expressed in Grain	Top Hit Descriptor	Homo sapiens golgi matrix protein GM130 (GOLGA2) mRNA, complete cds	bb72h12.y1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3047975 5' similar to gb:L08069 DNAJ PROTEIN HOMOLOG 2 (HUMAN);	bb72h12.y1 NIH_MGC_12 Homo saplens cDNA clone IMAGE:3047975 5' similar to gb:L08069 DNAJ PROTEIN HOMOLOG 2 (HUMAN);	Homo sapiens chromosome 21 segment HS21C048	Homo sapiens chromosome 21 segment HS21C046	EST389493 MAGE resequences, MAGO Homo saplens cDNA	EST389493 MAGE resequences, MAGO Homo saplens cDNA	Homo sapiens chromosome 21 segment HS210027	ar09d08.s1 Soares_testts_NHT Homo saplens cDNA clone IMAGE:1031151 3	Homo sapiens elF4E-like cap-binding protein (4EHP) mRNA	wo87501.x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2462281 3' similar to contains element	L1 repetitive element;	Homo sapiens chromosome 21 segment HS21 C046	Homo sapiens chromosome 21 segment HS21C046	Homo sapiens glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) (GOT2),	nuclear gene encoding mitochandrial protein, mrtvA	EST370215 MAGE resequences, MAGE Homo septens cDNA	EST370215 MAGE resequences, MAGE Homo sapiens cDNA	AU124387 NT2RM2 Homo sapiens cDNA clone NT2RM2002113 5	Homo sapiens anglopoletin 4 (ANG4) mRNA, partial cds	602123474F1 NIH_MGC_56 Hamo sapiens cDNA clone IMAGE:4280395 5	ozzgb03.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA cione IMAGE:16/6/17 3	H.sapiens dopamine receptor D5 pseudogene 1, partial cds	Homo sapiens ataxin 2-binding protein 1 (A2BP1), mRNA	QV1-HT0413-010200-059-h12 HT0413 Homo sapiens cDNA	Homo sapiens lymphocyte cytosolic protein 1 (L-plastin) (LCP1), mRNA	Homo sapiens lymphocyte cytosolic protein 1 (L-plastin) (LCP1), mRNA	Homo sapiens hypothetical protein SBBI87 (LOC57115), mRNA	AU132570 NT2RP4 Homo sapiens cDNA clone NT2RP4000109 5'	EST04286 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBUS88	EST04286 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBUS 08	602042882F1 NCI CGAP BING? Homo septens CONA Cigne INVAGE: 4 100330 3
ile Exon Prop	Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	Г	LNT		T_HUMAN	LN L	EST_HUMAN			T_HUMAN		TN			٦	\neg	EST_HUMAN			T_HUMAN			T_HUMAN		ΝT			П	╗	EST_HUMAN
Sing	Top Hit Acession No.	AF248953.1	BE206521.1	BE206521.1	AL163246.2	AL163246.2			AL163227.2	AA609940.1	4757701 NT		AI927030.1	AL163246.2	AL 163246.2		4504068 NT	AW958145.1	AW958145.1		AF113708.1	BF668537.1	A1078387.1	M77185.1	11431054 NT	AW606785.1	11434008 NT	11434008 NT	1.1423508 NT	AU132570.1	2.0E-64 T06397.1	2,0E-64 T06397.1	BF528114.1
	Most Similar (Top) Hit BLAST E Value	3.0E-64				3.0E-64 /	3.0E-64	3.0E-64	3.0E-64	2.0E-64	2.0E-64		2.0E-64	2.0E-64	2.0E-64			2.0E-64 /	2.0E-64	2.0E-84 /	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64
	Expression Signal	1.93	3.69	3.69	1.54	1.54	9.0	9.0	1.8	1.64	1.54		1.82	2.05	2.05		1.42	0.78	0.78	2.78	1.47	5.21	1.16	4.54	0.7	0.65	0.73	0.73	0.56	76.0	0.59	0.59	3.72
	ORF SEQ ID NO:	34199	34220	Į			35263	35264	37608	26484		l			27988	l _	28546	29157	29158	31655	31895			32378	ļ		١	L	L			ll	36592
	Exen SEQ ID NO:	21058	i	<u> </u>	L		L	L.	L		1	<u> </u>	15244	15248	L		15901	16519	1	İ	18925	ı	19263	ı	ı	20396	ı		1	1	ı	1	23353
	Probe SEQ ID NO:	8366	8393	8393	9327	9327	9414	9414	11691	1066	1377		2528	2533	2533		3137	3767	3767	5916	6148	6394	6497	6601	7077	7732	8567	8567	9038	9130	9879	9879	10662

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	wn81b06.x1 NCI_CGAP_Ut1 Homo saplens cDNA clone IMAGE:2452211 3'	wn81b06.x1 NCI_CGAP_Ut1 Hamo sapiens oDNA clone IMAGE:2452211 3'	PM2-SN0018-220300-002-e12 SN0018 Homo sapiens cDNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	CHR220101 Chromosome 22 exon Homo sapiens cDNA clone C22_132 6'	Homo saplens chromosome 21 unknown mRNA	au60c01.x1 Schnelder fetal brain 00004 Homo saplens cDNA clone IMAGE:2519138 3' similar to gb:L21696_cds1 PROTHYMOSIN ALPHA (HUMAN);contains element MSR1 repetitive element ;	Homo saplens synaptojanin 1 (SYNJ1), mRNA	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM3 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes,	complete cds; and L-type calcium channel a>	Homo sapiens TRIAD3 mRNA, partial cds	Homo sapiens TRIAD3 mRNA, partial cds	Homo saplens hypothetical protein FLJ11028 (FLJ11026), mRNA	zk53f08.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA done IMAGE:486567 3'	Homo sapiens chromosome 21 segment HS21C046	H.sapiens DNA for endogenous retroviral like element	H.saplens DNA for endogenous retroviral like element	QV4-BT0257-081199-017-e03 BT0257 Homo saplens cDNA	au58h07.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2519005 3' similar to SW:RL21_HUMAN P46778 60S RIBOSOMAL PROTEIN L21.;	QV2-BT0635-240400-162-c02 BT0635 Homo saplens cDNA	HSAAAEAWO TEST1, Human adult Testis tissue Homo sapiens cDNA clone cam test346 (b)	AV721898 HTB Homo sapiens cDNA clone HTBBZC06 5'	nj86d10.s1 NCI_CGAP_Pr11 Homo sapiens cDNA clone IMAGE:998379 similar to gb;K03002 60S RIBOSOMAL PROTEIN L32 (HUMAN);	nh37b07.s1 NCI_CGAP_Pr5 Homo sapiens cDNA clone IMAGE:954517	xe07b09.x1 NCI_CGAP_Co21 Homo sapiens cDNA clone IMAGE:2583545 3' sImilar to TR:Q63309 Q63309 LONG INTERSPERSED REPETITIVE DNA CONTAINING 7 ORF'S, ;contains L1.b2 L1 repetitive element ;	zw33b06.s1 Scares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773747 3	zw63b06.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA done IMAGE:773747 3'	qf18h05.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:1730426 3'
lle Exon Probes	Top Hit Database Source	EST_HUMAN wn	EST_HUMAN wn	EST_HUMAN PW	NT Ho	EST_HUMAN CH	Г		4507334 NT Ho						EST_HUMAN zk5		NT H.s	NT H.s	EST_HUMAN QV	est_HUMAN SW	EST_HUMAN QV		EST_HUMAN AV	EST_HUMAN RIB		xc0 EST_HUMAN LOI	EST_HUMAN zw	П	EST_HUMAN qf1
Sing	Top Hit Acession No.	Γ	2.0E-64 AI922911.1	1	37387	2.0E-64 H55162.1		Al928419.1	1.0E-64 4507334					1.0E-64 8922829 NT		1.0E-64 AL163246.2	9.0E-65 X89211.1		1.1	8.0E-65 A1929244.1	1		3.1		6.0E-65 AA503892.1	6.0E-65 AW083252.1		6.0E-65 AA427878.1	
	Most Similar (Top) Hit BLAST E Value	2.0E-64	2.0E-64	2.0E-64	2.0E-64	2.0E-64	1.0E-64	1.0E-64	1.0E-64		1.0E-64	1.0E-64	1.0E-64	1.0E-64	1.0E-64	1.0E-64	9.0E-65	9.0E-65	9.0E-65	8.0E-65	7.0E-85	7.0E-65	6.0E-65	8.0E-65	6.0E-65	6.0E-65	6.0E-65	6.0E-65	6.0E-65
	Expression Signal	5.97	5.97	1.76	1.55	4.85	2.94	10.45	0.79		5.74	1.27	1.27	0.79	1.07	1.81	1.53	1.53	10.43	10.87	2.01	1.27	3.59	4.73	0.62	23	3.46	3.46	0.81
	ORF SEQ ID NO:	36909	36910	37149	31114		25698	27214	28426		28912	28974	28975	29270	35816		27738	27739		37405	35921	37731	26454		32242	34481	34742	34743	34814
	Exon SEQ ID NO:	23656		23863	24562	24834	13060	14514	15776		16257	16327	16327	16631	22612		15000	15000	24123				13794	14652	19242	21337	21600	21600	21663
	Probe SEQ ID NO:	10981	10981	11198	12039	12468	251	1772	3010		3501	3572	3572	3881	9964	12012	2274	2274	11523	11493	10055	11807	1034	1915	6475	8645	8909	8909	8973

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Expn NO: CRF SEQ Signal ID NO: Expression Signal Signal No: Most Similar From ID NO: Top Hit Acession Signal No: Top Hit Acession No: Top Hit Acession Source 21663 34815 0.81 6.0E-65 BE567816.1 EST_HUMAN 23466 36707 3.82 6.0E-65 BE567816.1 EST_HUMAN 23644 36897 1.52 6.0E-65 BE567816.1 EST_HUMAN 24081 37392 1.6 6.0E-65 AL163210.2 NT 14080 26754 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 AC607848 INT 14080 26754 1.6 5.0E-65 A507848 INT 16012 28683 1.6 5.0E-65 A507848 INT 16012 28684 1.0 5.0E-65 A507848 INT 13502 26157 1.3 4.0E-65 A1266488.1 EST_HUMAN 13502 26157 1.3 4.0E-65 A1266488.1 EST_HUMAN 13502 27766								
21683 34815 0.81 6.0E-65 A1085314.1 EST_HUMAN 23466 36707 3.82 6.0E-65 BE567816.1 EST_HUMAN 23644 36897 1.52 6.0E-65 BE567816.1 EST_HUMAN 24081 37392 1.86 6.0E-65 A163210.2 NT 14080 26754 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 A503788.1 NT 16012 28683 1.6 5.0E-65 A507848 NT 16012 28684 1.6 5.0E-65 A507848 NT 16012 28684 1.6 5.0E-65 A507848 NT 13001 25642 2.02 4.0E-65 A120418.1 EST_HUMAN 13502 26156 1.37 4.0E-65 A12648.1 EST_HUMAN 13502 26157 1.37 4.0E-65 A126648.1 EST_HUMAN 15060 27777 0.91 4.0E-65		Ewn SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	当 ≠Ⅲ	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
23466 36707 3.82 6.0E-66 BE567816.1 EST_HUMAN 23644 36897 1.52 6.0E-65 BF340825.1 EST_HUMAN 24081 37392 1.86 6.0E-65 AL163210.2 NT 14080 26754 1.6 5.0E-65 AL163210.2 NT 14080 26755 1.6 5.0E-65 A507848 NT 16012 228684 1.6 5.0E-65 A507848 NT 16012 228684 1.6 5.0E-65 A507848 NT 1301 22662 1.01 5.0E-65 A507848 NT 1301 22664 1.01 5.0E-65 A507848 NT 1301 22642 2.02 4.0E-65 A120418.1 EST_HUMAN 13502 26157 1.37 4.0E-65 A126648.1 EST_HUMAN 13502 27796 0.91 4.0E-65 A26638 NT 1506 277797 0.91 4.0E-65 A26638 </td <td>8973</td> <td>21663</td> <td>34815</td> <td></td> <td>6.0E-65</td> <td>AI085314.1</td> <td>EST_HUMAN</td> <td>qf18h05.xf NCI_CGAP_Bm25 Homo sepiens cDNA clone IMAGE:1750425 3'</td>	8973	21663	34815		6.0E-65	AI085314.1	EST_HUMAN	qf18h05.xf NCI_CGAP_Bm25 Homo sepiens cDNA clone IMAGE:1750425 3'
25644 38897 1.52 6.0E-65 BF340825.1 EST_HUMAN 24081 37392 1.86 6.0E-65 AL16320.2 NT 14080 26754 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 7661951 NT 16012 28663 1.6 5.0E-65 4507348 NT 16012 28663 1.6 5.0E-65 4507348 NT 16012 28663 1.01 5.0E-65 A507348 NT 16012 28663 1.01 5.0E-65 A507348 NT 16012 28664 1.01 5.0E-65 A507348 NT 13502 26156 1.01 5.0E-65 A1267368 NT 13502 26156 1.01 5.0E-65 A1266488.1 EST_HUMAN 13502 27736 0.91 4.0E-65 A1266488.1	10783	23466	36707		6.0E-65	BE567816.1	EST_HUMAN	601340485F1 NIH_MGC_53 Homo sapiens cDNA done IMAGE:3682677 5
24081 37392 1.86 6.0E-65 AL163210.2 NT 14080 26754 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 7661951 NT 14080 26755 1.6 5.0E-65 7661951 NT 16012 28663 1.6 5.0E-65 4507848 NT 16012 28664 1.6 5.0E-65 4507848 NT 16012 28663 1.01 5.0E-65 4507848 NT 13001 25642 2.02 4.0E-65 AL120419.1 EST_HUMAN 13502 26156 1.37 4.0E-65 AL26748.1 EST_HUMAN 13502 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 <td< td=""><td>10968</td><td>23644</td><td>36897</td><td>1.52</td><td>6.0E-65</td><td>BF340825.1</td><td>EST_HUMAN</td><td>602037721F1 NCI_CGAP_Bm64 Homo sepiens cDNA clone IMAGE:4185677 5</td></td<>	10968	23644	36897	1.52	6.0E-65	BF340825.1	EST_HUMAN	602037721F1 NCI_CGAP_Bm64 Homo sepiens cDNA clone IMAGE:4185677 5
14080 26754 1.6 5.0E-65 AG61951 NT 14080 26755 1.6 5.0E-65 A6033768.1 NT 14080 26755 1.6 5.0E-65 A6033768.1 NT 16012 28643 1.6 5.0E-65 A507848 NT 16012 28644 1.6 5.0E-65 A507848 NT 13001 25642 2.02 4.0E-65 AL120418.1 EST_HUMAN 13502 26156 1.37 4.0E-65 AL26488.1 EST_HUMAN 13502 26156 1.37 4.0E-65 A266488.1 EST_HUMAN 13502 26156 1.37 4.0E-65 A506636 NT 13502 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16890 28221 1.03 4.0E-65 BE22	11480	24081	37392	1.86	6.0E-65	AL163210.2	N	Homo sapiens chromosome 21 segment HS21 CO10
14080 26755 1.6 5.0E-65 AB033768.1 NT 14881 27615 1.6 5.0E-65 AB033768.1 NT 16012 28663 1.6 5.0E-65 4507848 NT 16012 28664 1.01 5.0E-65 4507848 NT 23011 36226 1.01 5.0E-65 AF00868.1 NT 13502 26156 1.37 4.0E-65 AL120418.1 EST_HUMAN 13502 26157 1.37 4.0E-65 A1266488.1 EST_HUMAN 13502 26157 1.37 4.0E-65 A266688.1 EST_HUMAN 15060 27796 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.03 <	1331	14080	26754	1.6	5.0E-65	7661951	NT	Homo sapiens KIAA0156 gene product (KIAA0156), mKNA
14881 27615 1.6 5.0E-65 AB033768.1 NT 16012 28663 1.6 5.0E-65 AF009688.1 NT 16012 28663 1.6 5.0E-65 AF009688.1 NT 13001 25642 2.02 4.0E-65 AF009688.1 NT 13502 26156 1.37 4.0E-65 AL120419.1 EST_HUMAN 13502 26157 1.37 4.0E-65 AL266488.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266488.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266488.1 EST_HUMAN 15060 27796 0.91 4.0E-65 AL266488.1 EST_HUMAN 15060 277797 0.91 4.0E-65 BE221469.1 EST_HUMAN 16080 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16820 29321 1.03 4.0E-65 BE221469.1 EST_HUMAN 16820 29321 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 <td< td=""><td>1331</td><td>14080</td><td>26755</td><td></td><td>5.0E-65</td><td>7661951</td><td>NT</td><td>Homo sapiens KIAA0156 gene product (KIAA0156), mRNA</td></td<>	1331	14080	26755		5.0E-65	7661951	NT	Homo sapiens KIAA0156 gene product (KIAA0156), mRNA
16012 28663 ' 1.6 5.0E-65 4507848 NT 16012 28664 1.6 5.0E-65 AF009668.1 NT 23011 36226 1.01 5.0E-65 AF009668.1 NT 13001 25642 2.02 4.0E-65 AL120418.1 EST_HUMAN 13502 26156 1.37 4.0E-65 AL266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266468.1 EST_HUMAN 15060 27786 0.91 4.0E-65 AL266468.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.03 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.03 4.0E-65 BA093185.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BA093185.1 EST_HUMAN 18842 31804 4.0E-65 BA093185.	2151	14881	27615		5.0E-65	AB033768.1	NT	Homo sapiens hPAD-colony10 mRNA for peptidylarginine deiminase type I, complete cds
16012 28664 1.6 5.0E-65 4507848 NT 23011 36226 1.01 5.0E-65 AF009668.1 NT 13001 25642 2.02 4.0E-65 AL120418.1 EST_HUMAN 13502 26156 1.37 4.0E-65 AL266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 AL266468.1 EST_HUMAN 15060 27786 0.91 4.0E-65 AL26648.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.08 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BA093185.1 EST_HUMAN 18842 31804 4.0E-65 BA093185.1 EST_HUMAN 18842 31804 4.0E-65 BA093185.1 NT <td>3250</td> <td>16012</td> <td>28663</td> <td>1.6</td> <td>5.0E-65</td> <td>4507848</td> <td>M</td> <td>Homo sapiens ubiquitin specific protease 13 (isopeptidase 1-3) (USP13) mRNA</td>	3250	16012	28663	1.6	5.0E-65	4507848	M	Homo sapiens ubiquitin specific protease 13 (isopeptidase 1-3) (USP13) mRNA
23011 36226 1.01 5.0E-65 AF009668.1 NT 13001 25642 2.02 4.0E-65 AL120419.1 EST_HUMAN 13502 26156 1.37 4.0E-65 Al266468.1 EST_HUMAN 13502 26157 1.37 4.0E-65 Al266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 Al266468.1 EST_HUMAN 13816 26903 11.06 4.0E-65 Al266468.1 EST_HUMAN 15060 27786 0.91 4.0E-65 Al266468.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BW 993185.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BW 993185.1 NT 18842 31804 4.6 4.0E-65 BW 993185.1 NT 18676 32723 0.97 4.0E-65 BW 993303.1 <td>3250</td> <td>16012</td> <td>28664</td> <td>1.6</td> <td>5.0E-65</td> <td>4507848</td> <td>NT</td> <td>Homo saplens ublquitin specific protease 13 (isopeptidase 1-3) (USP13) mKNA</td>	3250	16012	28664	1.6	5.0E-65	4507848	NT	Homo saplens ublquitin specific protease 13 (isopeptidase 1-3) (USP13) mKNA
13502 2642 2.02 4.0E-65 AL120419.1 EST_HUMAN 13502 26156 1.37 4.0E-65 AL26648.1 EST_HUMAN 13602 26157 1.37 4.0E-65 AL26648.1 EST_HUMAN 13602 26475 1.38 4.0E-65 AL26648.1 EST_HUMAN 13604 27784 1.38 4.0E-65 BE22146.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 28321 1.08 4.0E-65 BE221469.1 EST_HUMAN 16880 28321 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BW993185.1 NT 18842 31804 4.0 4.0E-65 BW993185.1 NT 19709 32765 0.97 4.0E-65 BW993189.1	10364	23011	36226		5.0E-65	AF009668.1	IN	Multiple sclerosis associated retrovirus polyprotein (pol) mRNA, partial cds
13502 26156 1.37 4.0E-65 Al266468.1 EST_HUMAN 13502 26157 1.37 4.0E-65 Al266468.1 EST_HUMAN 13814 26475 1.38 4.0E-65 Al266468.1 EST_HUMAN 15060 27786 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.08 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 BE221469.1 EST_HUMAN 17842 30450 1.03 4.0E-65 Aw993185.1 NT 17842 30460 1.03 4.0E-65 Aw993185.1 NT 18842 31805 4.0 4.0E-65 Aw993185.1 NT 19709 32765 0.97 4.0E-65 My99318.1 NT 20124 33216 0.97 4.0E-65 My99318.1 NT 20437 33559 1.86 4.0E-65 U40372.1 NT	188	13001	25642	2.02		AL120419.1	EST_HUMAN	DKFZp761G108_r1 761 (synonym: hamy2) Homo sapiens cDNA clone DKrZp761G108 5
13502 26157 1.37 4.0E-85 Al266468.1 EST_HUMAN 13814 26475 1.38 4.0E-85 4826735 NT 14216 26903 11.06 4.0E-65 4506636 NT 15060 27786 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27787 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 28321 1.08 4.0E-65 AV963185.1 EST_HUMAN 17842 30450 1.03 4.0E-65 AV963185.1 EST_HUMAN 17842 30460 1.03 4.0E-65 AV963185.1 EST_HUMAN 17842 30460 1.03 4.0E-65 AV0633093.1 NT 1842 31805 4.0 4.0E-65 AY008372.1 NT 19709 32765 0.97 4.0E-65 AY008372.1 NT 20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33559 1.86 4	728	13502			4.0E-65	AI266468.1	EST_HUMAN	qm46e01.x1 Soeres_placenta_8tx9weeks_ZNbHP8tx9W Homo sapiens cDNA clone IMAGE:1891800 3'
13814 26475 1.38 4.0E-65 4826735 INT 14216 26903 11.06 4.0E-65 4506636 INT 15060 27796 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 16890 28321 1.08 4.0E-65 AW993185.1 EST_HUMAN 17842 30459 1.03 4.0E-65 AW993185.1 EST_HUMAN 17842 30460 1.03 4.0E-65 AW993185.1 EST_HUMAN 18842 31804 4.6 4.0E-65 AR933185.0 INT 19676 3273 0.55 4.0E-65 AR933093.1 INT 19709 32765 0.97 4.0E-65 AY008372.1 INT 20124 33216 0.97 4.0E-65 U40372.1 INT 20437 33558 1.86 4.0E-65 U40372.1 INT 20437 33559 1.86 4.0E-65 E433766 INT </td <td>728</td> <td>13502</td> <td></td> <td></td> <td>4.0E-85</td> <td>A1266468.1</td> <td>EST_HUMAN</td> <td>qm46e01.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:1891800 3'</td>	728	13502			4.0E-85	A1266468.1	EST_HUMAN	qm46e01.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:1891800 3'
14216 26903 11.06 4.0E-65 4506636 NT 15060 27796 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 16890 28321 1.08 4.0E-65 AW993185.1 EST_HUMAN 17842 30459 1.03 4.0E-65 AW993185.1 EST_HUMAN 17842 30460 1.03 4.0E-65 AW993185.1 EST_HUMAN 18842 31804 4.6 4.0E-65 AB032093.1 NT 19676 32723 0.55 4.0E-65 AY008372.1 NT 20124 32215 0.97 4.0E-65 AY008372.1 NT 20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33558 1.86 4.0E-65 U40372.1 NT 20437 33559 1.86 4.0E-65 E433766 NT 20437 34888 0.68 4.0E-65 B4	1058	13814			4.0E-65		TN	Homo sapiens fragile X mental retardation, autosomal homolog 1 (FXR1), mRNA
15060 27796 0.91 4.0E-65 BE221469.1 EST_HUMAN 15060 27797 0.91 4.0E-65 BE221469.1 EST_HUMAN 16880 29321 1.08 4.0E-65 AW993185.1 EST_HUMAN 17842 30459 1.03 4.0E-65 8055289 NT 17842 30460 1.03 4.0E-65 8055289 NT 18842 31804 4.6 4.0E-65 AB05209 NT 18842 31805 4.6 4.0E-65 AB033093.1 NT 19676 32723 0.55 4.0E-65 AY008372.1 NT 19677 32755 0.97 4.0E-65 AY008372.1 NT 20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33559 1.86 4.0E-65 U40372.1 NT 20437 33559 1.86 4.0E-65 E433766 NT 20437 33559 1.86 4.0E-65 E433766	1469	14216			4.0E-65		IN	Homo sapiens ribosomal protein L34 (RPL34) mRNA
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17842 30456 1.03 4.0E-65 9055269 NT 17842 30460 1.03 4.0E-65 9055269 NT 18842 31804 4.6 4.0E-65 AB033093.1 NT 18842 31805 4.6 4.0E-65 AR033093.1 NT 19676 32723 0.55 4.0E-65 AY008372.1 NT 19807 32873 2.52 4.0E-65 M19879.1 NT 20124 33215 0.97 4.0E-65 U40372.1 NT 20437 33216 0.97 4.0E-65 U40372.1 NT 20437 33559 1.86 4.0E-65 E463766 NT NT 20437 33559 1.86 4.0E-65 E463766 NT NT 20437 34888 0.63 4.0E-65 H1429127 NT	3930	16680	29321		4.0E-65	AW9931	EST_HUMAN	RC2-BN0033-160200-013-e03 BN0033 Homo sapiens cDNA
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19676 32723 0.55 4.0E-65 AY008372.1 NT 19709 32765 0.97 4.0E-65 M19879.1 NT 19807 32873 2.52 4.0E-65 M1545780 NT 20124 33215 0.97 4.0E-65 U40372.1 NT 20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33558 1.86 4.0E-65 U40372.1 NT 20437 33559 1.86 4.0E-65 E453765 NT 21734 34888 0.63 4.0E-65 E453765 NT	6063	18842				AB033093.1	Ę	Homo sapiens mRNA for KIAA1267 protein, partial cds
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19807 32873 2.52 4.0E-65 11545780 NT 20124 33215 0.97 4.0E-65 U40372.1 NT 20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33558 1.86 4.0E-65 S453765 NT 20437 33559 1.86 4.0E-65 S453765 NT 21734 34888 0.63 4.0E-65 T429127 NT	7017	19709			4.0E-65	M19879.1	۲N	Human clabindin 27 gene, exons 10 and 11, and L1 and Au repeats
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20124 33216 0.97 4.0E-65 U40372.1 NT 20437 33558 1.86 4.0E-65 5453765 NT 20437 33559 1.86 4.0E-65 5453765 NT 21734 34888 0.63 4.0E-65 1429127 NT	7448	20124	L			U40372.1	TN	Human 3',5' cyclic nucleotide phosphodiesterase (HSPDE1C3A) mRNA, partial cds
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20437 33559 1.86 4.0E-65 5453765 NT 21734 34888 0.63 4.0E-65 11429127 NT	7741	20437					LZ	Homo saplens nel (chicken)-like 2 (NELL2), mRNA
21734 34888 0.63 4.0E-65 11429127 NT	7741	20437					된	Hamo sapiens nel (chicken)-like 2 (NELL2), mRNA
	9044	21734	L		4.05		LΝ	Homo sapiens Janus khase 2 (a protein tyrosine kinase) (JAK2), mKNA
23126 2.04 4.0E-65 AJ277546.2 NT	10480	23126		2.04	4.0E	AJ277546.2	TV LV	Homo saplens WEE1 gene for protein kinase and partial ZNF143 gene for zinc finger transcription factor

Page 340 of 536 Table 4 Sindle Exon Probes Expressed In I

Single Exon Probes Expressed in Brain	ORF SEQ Expression (Top) Hit Top Hit Acession (Top) Hit Top Hit Acession (Top) Hit Descriptor Signal BLASTE No. Source	36801 1.97 4.0E-65 AV738764.1 EST_HUMAN AV738784 CB Homo sapiens cDNA clone CBCCBE06 6"	36982 3.68 4.0E-65 AF119846.1 NT Homo saplens PRO1474 mRNA, complete cds	29475 1.46 4.0E-86 4826736 NT Homo sapiens fragile X mental retardation, autosomal homolog 1 (FXR1), mRNA	3.8 3.0E-65 X78932.1 NT H.saplens HZF9 mRNA for zinc finger protein	26984 0.91 3.0E-65 4504626 NT Homo sepiens immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	ov23703.s1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:1638173 3' similar to contains element 27271 0 83 3 0E-85 A1000892.1 FST HIMAN MSR1 repetitive element:	0.75 3.0E-65 4604960 NT	ov23f03.s1 Scares_testis_NHT Homo explens cDNA clone IMAGE:1638173 3' similar to contains element 29101 0.99 3.0E-66 A1000692.1 EST HUMAN MSR1 repetitive element;	1.91 3.0E-65 6912385.NT	3.0E-65 BE787366.1 EST HUMAN	3.0E-66 AA430006.1 EST_HUMAN	35 BF680294.1 EST_HUMAN	35 BE263373.1 EST_HUMAN	32.07 2.0E-65 BF676922.1 EST_HUMAN	1.06 2.0E-65 AK024463.1 NT	34563 1.06 2.0E-86 AK024463.1 NT Homo septiens mRNA for FLJ00056 protein, partial ods	2	2.0E-65 AA307904.1 EST_HUMAN	35 BF246086.1 EST_HUMAN	1.0E-65 BF126644.1 EST_HUMAN	1.44 1.0E-65 7867496[NT	2749B 1.29 1.0E-66 AB040946.1 NT Homo sepiens mRNA for KIAA1513 protein, partial eds	1.0E-66 BE466681.1 EST_HUMAN	1.0E-66 4504082 NT	2.47 1.0E-65 4504082 NT	1.0E-65 AW029340.1	2.01 1.0E-65 AW029340.1 [EST_HUMAN	2.04 1.0E-85 AW820481.1 EST HUMAN	33978 2.04 1.0E-65 AW820481.1 EST_HUMAN QV2-ST0298-140200-042-f12 ST0298 Homo septens dDNA
	Exen ORF S SEQ ID ID NO:	23664 36	23712 36	13814 26	15522	14297 26	14556 27		16462 29		22617 35		16157 28	19210			21436 34	23248 36	24515	25156	12915			16124 28		16728 29				20846 33
	Probe SEQ ID SE NO:	10874	11041	L	1210	1551	1816	1	3709	L	L			6442	1		8744			12431			2033	L		3980	4183	1		8152

Page 341 of 536 Table 4 Exon Probes Expressed in Brain

L XIII OCCUPATION OF THE PROPERTY OF THE PROPE	Too Hit Descriptor	10p.1	601 566124F1 NIH_MGC_21 Homo saplens cDNA closs IMAGE:3841012 5	601566124F1 Nin	AU141283 THYRO1 Homo saplens cDNA clone IMAGE:4283313 5	A0141200 11H MGC_56 Homo saplens COUNTY 2004714 5	602120A331 VIII Homo sapiens cDNA clone N. 2.0	AU (2804) NT2RP2 Homo septens cDNA doller (TPR1), mRNA	AU 1250- Homo saplens Inositol 1,4,8-triphosphate receptor, MRNA Homo saplens Inositol 1,4,8-triphosphate (KIAA0656), mRNA	Homo sepiens KIAAU000 Benor Land sapiens cDNA clone Ilway Carrier element:	dd56a02.x1 Soares tests Int I the MER19.t1 MER19 appearate Commission of the Management of the Meritan State of th				T	Homo sapiens ribosomal protein L. a (**) Homo sapiens ribosomal population (**) Homo sapiens ribosomal population (**)	Т	\top		1 1	Homo sapiens sulfotransferase-related process	Novel human gene mapping to chomosome 22	Novel human gene mapping to the part homolog (PORI) mRNA	Homo saplens 265 processoriated pad1 homotog (POLI)	Homo sapiens 200 F. Home sapiens 200 F. Home MAGE:767048 5	Human transposed Time S1 Homo sapiens clura contractions of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations of the S1 Homo sapiens clurations clura	۲	7		Т	_ 1	
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·	+	Most Similar Top (Top) Hit		1.0E-65 BE	1.0E-65 BE	1.0E-65 AU	1.0E-65 AL	1.08-89	1.0E-65 AU129040.1	1 OF-85	101		1.0E-65/	1.05-83	1000	4 OF -65	100	1.05-50 1.0F-65		1.0E-65	1.05-05		L	١.	١	L	\square		Ц	,	1.15	1.15 6.00
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Page 342 of 536 Table 4 Single Exon Probes Expressed in Brain

	Top Hit Descriptor	wn57h07.x1 NCI_CGAP_Lu19 Homo seplens cDNA clone IMAGE:2449697 3' similar to WP:F16G9.4A CE18695 ;	PM2-HT0604-030300-001-b06 HT0604 Homo saplens cDNA	protein 131	RC4-BT0311-141199-011-h06 B10311 Home sapiens culvA	601681692F1 NIH MGC 9 Homo sapiens con A cicle IMAGE: 393 179 1 V	60168159271 NIH MGC 9 Homo septens conta civila invade. 393 i 70 i 6	Homo sapiens (nyroid normone receptor birtaing process), mission	Mus musculus fragile X mental retargation syndiome 1 normaly (1 nn 1), mayor	RC1-NN0063-100500-022-a02 NN0063 Homo sapiens culve	ous refroviral like element	Homo sapiens gem-line DNA upstream of Jkappa locus	is, complete genome	Homo saptens methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenytetranydrororate ovclohydrolase (MTHFD2), mRNA	QV1-DT0069-110200-067-910 DT0069 Homo saplens cDNA	EST377646 MAGE resequences. MAGI Homo saplens cDNA		Homo sapiens cAMP-regulated guanine nuclectide exchange factor I (cAMP-GEFI) mRNA, complete cds	Homo sapiens metrylene tetrahydrotolate denydrogenase (NAD+ dependent), metrentyleu arrydrotolate cyclohydrolase (MTHFD2), mRNA	Homo sapiens hypothetical protein FLJ20116 (FLJ20116), mRNA	Js pHE.1 (ERV9)	UI-H-BW1-amr-a-10-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3070747 3	Homo sapiens mRNA for KIAA0998 protein, partial cds	Homo septiens solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member o (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA	Homo sepiens solute carrier family 26 (mitochondrial carrier; adenine nucleotide translocator), member 6 (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA	yz27g12.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:284326 6' similar to	SW:H2B1_TIGCA P36068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;	yzzgg12.r1 Soares_multiple_scienceis_znbrintsir homo sapiens cultura ciute invocezgg25.g ciining SW:HzB1_TIGCA P35068 HISTONE H2B.1/HzB.2. [2] PIR:B56612;
Single Excit Flobes Explessed in Didin		wn57h07.x1 NCI_CGAP_Lu CE18595;	PM2-HT0604-030300-001-b	H.sapiens mRNA for ribosomal protein L31	RC4-BT0311-141199-011-h	601681592F1 NIH_MGC_9	601681592F1 NIH MGC_9	Homo saplens tnyrold normo	Mus musculus fragile X mer	RC1-NN0063-100500-022-6	H.saplens DNA for endogenous refroviral like element	Homo sapiens germ-line DN	Human endogenous retrovirus, complete genome	Homo saplens methylene tetrahydr cyclohydrolase (MTHFD2), mRNA	Т	Γ		Homo sapiens cAMP-regula	Homo sapiens methylene tetrehydr cyclohydrolase (MTHFD2), mRNA	Homo sapiens hypothetical	Human endogenous retrovirus pHE.1 (ERV9)	П		Homo saplens solute carrie (SLC25A5), nuclear gene e	Homo sapiens solute carrie (SLC25A5), nuclear gene e	yz27g12.r1 Soares_multiple		$\neg \neg$
I EXOII LIO	Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT .	EST HUMAN	EST_HUMAN	EST HUMAN	NT	L	EST_HUMAN	NT	NT	NT	L _Z	EST HUMAN	FOT UNIVERSE	NEWOL I CH	N	F	ΙΝ	Į.	EST_HUMAN	N	۲	Į		EST HUMAN	EST HUMAN
SIIC	Top Hit Acession No.	66 Al924653.1		66 X69181.1	66 BE064410.1	66 BE898644.1	G BE898644.1	11420557 NT	6679816 NT	66 AW897798.1	66 X89211.1	4.0E-66 AJ223364.1	9635487 NT	TN 528643	AW939	A 071 700111	-66 AW865473.1	-66 U78168.1	TN 629643		X57147	-66 BF507493.1	-66 AB023215.1	4502098 NT			-66 N55323.1	3.0E-66 N66323.1
	Most Similar (Top) Hit BLAST E Value	8.0E-66	8.0E-66	6.0E-66	5.0E-66	5.0E-68	5.0E-66	5.0E-66	4.0E-66	4.0E-66	4.0E-66	4.0E-66	4.0E-66	89 110 1	4.0E-00		4.0E-66	4.0E-66	4 OF 86	4 0E-66	4 0F-86	4.0E-66	4.0E-66	E S		30.5	3.0€	
	Expression Signal	1.15	0.48	3.14	1.45	0.74	0.74	16.11	0.98	1.14	1.83	3.02	10.89	i.	1 15	2	4.83	7.83	5	E 63	0.73	1.97	1.28	'		08.01	1.18	1.16
	ORF SEQ ID NO:	29700		37062	28769	30382	30383	35028	26206	27170	27744						30561	32779	_	21.00	1		L		1	92897	27429] .
	SE SON NO:	17071	21026	1	ı	ł	L		13545	14471	1	L	L	<u> </u>		70+01	17926	19723	1	1	7007	1	1	ļ		14154	14711	1 1
	Probe SEQ ID NO:	4332	8333	11108	1346	5046	5046	9194	733	1729	2278	2477	4733	3	5463	/000	6757	7031		RZQ/	0/8/	10004	44284	3	J45	1407	1975	1975

Page 343 of 536 Table 4 Single Exon Probes Expressed in Brain

I Top Hit Database Source		727/912.r1 Soares_multiple_sclerosis_2NbHMSP Homo sepiens cDNA clone IMAGE:284326 6' similar to SW:H2B1_T1GCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612:	Homo sapiens TGF(beta)-induced transcription factor 2 (TGIF2), mRNA	Homo saplens KIAA0649 gene product (KIAA0649), mRNA	Homo sapiens mRNA for KIAA0892 protein, partial cds	Homo sapiens protein kinase C beta-II type (PRKCB1) mRNA, complete cds	Homo sapiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1), mRNA	Homo sapiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1), mRNA	Homo sapiens mRNA for FLJ00045 protein, partial cds	Homo sapiens KIAA0433 protein (KIAA0433), mRNA	Homo sapiens protocadherin beta 1 (PCDH-beta1), mRNA	Homo sapiens molyodenum cofactor biosysthesis protein E (MCBPE) mRNA, complete ods	Homo saplens protein phosphatase 2, regulatory subunit B (1856), alpha isoform (PPP2R5A) mRNA	Homo sepiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and pertial cds, alternatively spliced	Homo saciens Misshapen/NiK-related kinase (MinK) mRNA	Homo sapiens Misshapen/NIK-related Knase (MINK), mRNA	Homo sapiens origin recognition complex, subunit 5 (yeast homolog)-like (ORC5L) mRNA, and translated products	Homo sepiens origin recognition complex, subunit 5 (yeast homolog)-like (ORCSL) mRNA, and translated	products	Homo sapiens chromosome 21 segment HS21C101	Hamo sapiens hypothetical protein FLJ20309 (FLJ20309), mRNA	Novel numen gene mapping to chomosome 1	Homo saplens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds	Homo sapiens HLA-B gene for human leucocyte antigen B	Homo sapiens HLA-B gene for human leucocyte antigen B	EST380830 MAGE resequences, MAGJ Homo sapiens cDNA	EST380930 MAGE resequences, MAGJ Homo sapiens cDNA	yy59c02.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:277826 6	Homo sapiens G-2 and S-phase expressed 1 (GTSE1), mRNA	601508376F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909931 5'	AV717817 DCB Hamo sapiens aDNA clane DCBADC07 6'
Top Hit Database Source		X EST_HUMAN S			I									H IS									Ĭ					EST_HUMAN yy		Г	EST HUMAN A
Top Hit Acessian No.		-68 N55323.1	1141880	7662223 NT	66 AB020699.1	66 M13975.1	11417946 NT	11417946 NT	Γ	-66 11417118 NT	7019480 NT	66 AF155659.1	5453949 NT	68 AF223391.1	57334	7657334 NT	4505524 NT		2524	-66 AL163301.2	2290						14.1	36 N45480.1 E	11418318 NT		36 AV717817.1 E
Most Similar (Top) Hit BLAST E		3.0E-88	3.0E-68	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66	3.0E-66 /	2.0E-66	2.0E-68	2.0E-66	1	2.05-00	2.0E-55	2.0E-88	2.0E-00 A	2.0E-68 A	2.0E-66 A	2.0E-66 A	2.0E-66 A	2.0E-66 A	2.0E-66 N	2.0E-66	1.0E-66 B	1.0E-68 A
Expression Signal	1	1.16	3.54	6.3	1.14	0.73	1.92	1.92	0.62	0.52	0.8	0.97	6.16	1.57	2.15	2.15	1.76	, .	1./0	207	0.77	00.0	200	9.48	9.48	1.3	1.3	2.26	2.37	1.15	1.36
ORF SEQ ID NO:		27431	28156	28520	30870		31397		35275	35459	35826	36287	37406	37730	25505	25506	25437	20730	22438	21212	75850	20447	71437	29973	29974	31439	31440	34585			28309
Exan SEQ ID NO:		14711	15418				18479		1			23066	24095	24396	12879	12879	12824	10001	47071	000	0070	2000	8 2	1/342	1/342	18518	18518	21438	25370	14422	15662
Probe SEQ ID NO:		1975	2711	3115	5380	5490	5686	5686	9425	9619	9973	10420	11494	11806	ଞ	. 20	413	-	2 0	8 0 8	2777	7707		į į	460/	5726	5728	8746	12329	1678	2885

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Table 4
On Probes Expressed

SEC D Cope Econoce Cope Econoce					_	_	_																										
SEC ID ORF SEC Expression Top Hit Acession	obes Expressed in Brain	Top Hit Descriptor	ba72g06,y1 NIH_MGC_20 Homo septens cDNA clone IMAGE:2806976 6' similar to TR:094892 О94892	historias Pro I EIN. be72g05.y1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2905978 5' similar to TR: Courses Courses	IKIAA0798 PROTEIN.;	Homo sapiens hypothetical protein dJ462023.2 (DJ462023.2), mRNA	Homo sapiens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	nomo sapiens KRAB zinc finger protein ZFQR mRNA, complete ods	round sapiens developmentally regulated GTP-binding protein 1 (DRG1), mRNA	Lastigut. St. Sogres, testis, NHT Homo saplens cDNA clone IMAGE:746392.3/	No. 2 Line Andrews Chromosome 21 segment HS21C100	Nove nurtain gene mapping to chomosome 13	6018/6351F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4091893 5	Homo saplens mRNA for NADPH-cytochrome P-450 reductase, complete cds	Hamo saplens mRNA for NADPH-cytochrome P-450 reductase, complete cds	DNF 22/01A229_r1 761 (synonym: hamy2) Homo saplens cDNA clone DKFZp761A229 6	ES I 3650 Embryo, 9 week Homo sapiens cDNA 5' end similar to similar to cerebellin	ES I 30530 Embryo, 9 week Hamo saplens CDNA 5' end similar to similar to carebellin	RC4-B10508-170100-011-c07 BT0568 Homo saplens cDNA	N.724660 1125 170100-011-007 BT0568 Homo sapiens cDNA	AV/31333 H I P Homo saplens cDNA clone HTFARD03 6/	ANA RESPONDE THE TOTAL OF THE SUB Home sapiens cDNA clone IMAGE: 2727283 3'	US 14552527 I NIT MCC_66 Home saplens cDNA clone IMAGE:3858978 51	SOLATETEDER NITH 1999 Protein (KIAA0985), mRNA	DM2 TN0103 0.0000 000 000 000 000 000 000 000 00	The control of the co	Home saprens myond autoantigen 70kD (Ku antigen) (G22P1), mRNA	inding separatis gamma-glutamyfitansferase 1 (GGT1), mRNA	Homo sapiens emyloid beta (A4) precursor protein (protesse nextn-II. Alzheimer disesse) (ADD)	2190b04.s1 Soares_fetal_liver_splean_1NFLS_S1 Homo sapiens cDNA clone IMAGE-446048.9	WD65c12.X1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2310550 3'	20144938871 NIH_MGC_66 Homo sapiens cDNA clone IMAGE;3852254 6; 2482h10.r1 Stratagene hNT neuron (#837233) Homo sapiens cDNA clone MACE;04446	SW:SAV_SULAC 007590 SAV PROTEIN :
E Exon ORF SEQ ID NO: Expression Signal Most Similar Top Hit Acess Signal BLAST E No. No. 7 14614 27325 1.36 2.0E-67 BE303037.1 14222 1 14614 27325 1.36 2.0E-67 BE303037.1 14223 1 14614 27325 1.36 2.0E-67 AF309561.1 14224 1 14615 27702 1.3 2.0E-67 AF309561.1 14224 2 16153 277845 1.09 2.0E-67 AF309561.1 14224 2 16163 277845 1.09 2.0E-67 AF309561.1 14224 2 16176 27846 1.09 2.0E-67 AL1633002.2 1462474.1 2 16176 27847 1.2 2.0E-67 AL163066.1 2.0E-67 AL16306.1 2 16176 27847 3.13 2.0E-67 AL16306.1 2.0E-67 AL16306.1 2 18878 3.1956 2.46 2.0E-67 AL16306.1 2.0E-67 AL16306.1 2 18878 3.1957 2.0E-67 AL16306.1 2.0E-67 AL16306.1 2.0E-67 AL16306.1 2 1888 3.7249 0.82 2.0E-67 AL16306.1 2.0E-67 AL16306.1 <td>ngle Exon Pr</td> <td></td> <td>FOT HISAAN</td> <td>NUMBER OF THE PARTY OF THE PART</td> <td>EN TOWAR</td> <td>1</td> <td>- N</td> <td>1</td> <td>TOT TOTAL</td> <td>NAME OF PARTY</td> <td>Ė</td> <td>ECT UNIVERS</td> <td>ES - HOMAN</td> <td>FIA</td> <td>ECT UITABL</td> <td>EST HOMAN</td> <td>TOT TOTAL</td> <td>EST LINANI</td> <td>EST HOWAIN</td> <td>EST CLIMAN</td> <td>EST LIMAN</td> <td>TST HIMM</td> <td>1000</td> <td>TST HIMAN</td> <td>ST HIMAN</td> <td></td> <td> L</td> <td></td> <td></td> <td>HOMAN</td> <td>T</td> <td>\neg</td> <td></td>	ngle Exon Pr		FOT HISAAN	NUMBER OF THE PARTY OF THE PART	EN TOWAR	1	- N	1	TOT TOTAL	NAME OF PARTY	Ė	ECT UNIVERS	ES - HOMAN	FIA	ECT UITABL	EST HOMAN	TOT TOTAL	EST LINANI	EST HOWAIN	EST CLIMAN	EST LIMAN	TST HIMM	1000	TST HIMAN	ST HIMAN		L			HOMAN	T	\neg	
Dec Exon ORF SEQ Expression Top) Signal Not	ĪS		BE303037.1	RE303037 4	1140004	11422040	AF309564 4	4768705	AA625755 1	Al 163300 2	AL049784 1	3F240758 1	AR084783 4		Γ	A334600 1	A334600 1	T	T		1		36448			8180	11417877 N		8		T.		
Column C		Most Similar (Top) Hit BLAST E Value	2.0E-67		2.0E-67	2.0F-67	2.0E-67	2.0E-67	2.0E-67	2.0E-67	2.0E-67	2.0E-67	2.0F-67	2 0F-67	2.0E-67	2.0E-67	2.0E-67	2.0E-67	2.0E-67	2.0E-87.A	2.0E-67 A	2.0E-67 B	2.0E-67	2.0E-67 B			2.0E-67	10 10		1 0F 87 A1	8.0F-68 RF	0 0 0 0	0.05-20174
E E E E E E E E E E E E E E E E E E E		Expression Signal	1.36	1.36	1.3	1.3	1.09	1.28	3.8	3.03	9.0	5.54	2.46	2.46	0.76	0.82	0.82	1.21	1.21	0.83	0.97	3.72	.4.67	2.11	1.86	1.36	205	0 34	101	8.58	. 2.13	4 08	200
8 C C C C C C C C C C C C C C C C C C C												31774	31956	31957	32315	34290	34291	34724	34725	35246	36447	36899		37145	36433	30819	31018	25695	26114	37770	27636	29238	
Probe SEQ ID NO: 1877 1877 2236 2236 2236 2236 2236 6203 6203 6203					ı	l		_ [- 1	ı	- 1		Ι.	į l	19310	21148	21148	21686	21586	22076	22261	23646	25433	23868	23201	2523	24874	13055	13467	24429	14903	16601	
		Probe SEQ ID NO:	1877	1877	2235	2236	2384	2432	3460	3884	5981	9034	6203	6203	6546	8466	8456	8895	8895	9466	9608	10970	10988	1193	11434	12235	12527	246	692	11846	2174		1

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Table 4
Single Exon Probes Expressed in Brain

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	HSDN 6778 FM ST FM	The First of History Replies CDNA clone \$3000023D09	Unicetalias longicaudatus mRNA for EF-1 alpha, complete cds	from suprems gene for activity receptor type IIB, complete cds	1938g04.s1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:34896 3'	021400141 Milliamor_go figure sapiens cone cione IMAGE:3862034 5	FORMIN 4 // IMB DEFORMITY DEOCEMAN	VZZRAOZ 11 Spares militale coleccio April 1165 II	OVO-BT0074-130999-014-014 BT0074 Home Series CENA Clone IMAGE:289165 6	Homo sabiens meningiome (Algerin/ed February Februa	QV4-ST0234-181199-037-105 ST0234 Home condens - DNA	Homo sabiens mBNA for KIAA0572 models	Homo saplens mRNA for KIA40577 makein complete ads	601177002F1 NIH MGC 17 Home spallene and all the property of t	847612 st Sognes NET TORO ST Demonstrate This September 1997	Homo saniens cell reconnition malecule Control (VIA Acces)	Homo carlana cartir (C. 140)	2w7440213 Sprage factor MLT Users	Homo sanlens bineshadischemen 28 (DDE26) - ENTA	Homo saplens phosphodiactores 78 (PDE/D), INRNA	Homo sapiens MIF2 suppressor (HSM73) "PNA con-122 J.	Homo saplens myosin IC (MYO1C), mRNA	Homo sapiens Interleukin-7 receptor precursor (II 7R) dens expres 7 and 8 and complete of	Human protein kinase C substrate 80K-H (PRKCSH) gene, even 4-5	Human protein kinase C substrate 80K-H (PRKCSH) gave even 4-5	Homo saplens CGI-76 protein (LOC51632) mRNA	Homo saplens CGI-76 protein (LOCS1832) mRNA	Homo sapiens meningloma (disrupted in balanced franshocation) 1 (MM11)	Homo saplens low density lipoprotein-related protein 2 (I RB2) mBNA	Homo sapiens pre-B-cell convenhancing factor (PBEE) with A	Homo sapiens pre-B-cell colony-enhancing factor (PBEE) mBNA	Homo saplens 26S protessome-associated pad1 homolog (PDH1) "BNA	Homo sapiens 28S proteasome associated pad1 homolog (POH1) mRNA	Homo saplens EphA4 (EPHA4) mRNA
jle Exon Prok	Top Hit Database Source	FOT HIMANI	NICINICIA I	- LN	TOT LINESAL	EST HUMAN	EST LIMAN	SWISSPROT	Т	Т	Т	T HUMAN	Г		T HUMAN	Т			T HUMAN							Ę									
Sing	Top Hit Acession No.	68 F28784 1		ļ	Ī	Ţ.	T			=	05222			Π	Γ	Γ	32340	11438718 NT		688	1.0E-68 11418869INT		33277	9.1			11418431 NT	11418431 NT	450522 NT	11430460 NT	5031976 NT	5031976 NT	5031980 NT	5031980 NT	4758279 NT
	Most Similar (Top) Hit BLAST E Value	3.05-68	200	2.0E-88	2 OF AB	2.0E-68	2.0E-68	2.0E-88	2.0E-68	2.0E-88	1.0E-68	1.0E-68	1.0E-68	1.0E-68	1.0E-68	1.0E-68	1.0E-68	1.0E-88	1.0E-68 A	1.0E-68	1.0E-68	1.0E-68 L76416.1	1.0E-88	1.0E-88 A	1.0E-68 U50319.1	1.0E-68 U50319.1	1.0E-68	1.0E-68	1.0E-68	1.0E-68	9.0E-89	9.0E-89	9.0E-89	9.0E-69	9.0E-69
	Expression Signal	1,23	15 31	1.38	0 54	5.38	0.73	0.63	0.75	1.66	Ψ-	13	1.03	1.03	0.95	0.92	1.37	1	9.0	1.85	1.85	3.5	1.71	1.62	1.28	1.26	1.48	1.48	2.66	2.11	1.16	1.16	1.41	1.41	1.15
	ORF SEQ ID NO:	36261		30004		32462	L		36376	37160	25541	25737	27715	27716	29373	30302	30674	33350	36235	36685	36686	36741	37098	37176	37223	37224	37584	37585	25541	30712	25460	25481	28426	28427	27740
	SEQ ID NO:	23045	L	1	19520	L	19954	ı	23151		12903	13095	14977	14977	16739	17694	18045	20245	23019	23441	23441	23502	23816	23880	23831	23831	24261	24261	12903	25322	12847	12847	13766	13766	15001
	Probe SEO ID NO:	10389	2865	4633	6776	සමය	7270	8848	10505	11210	*	88	2249	2249	3991	4969	6239	7676	10373	10756	10756	10819	1148	11226	11270	11270	11666	11986	12511	12719	9	9	100	100	2270

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	f	Τ	T	T	Т	Т	T	7	Т	Т	T	Т	Т	Т	1	Т	Т	Τ	Т	Τ-	Т	T	Т	Т	7	Т	Т
Oligio Lyon I Tobas Lypressed III Dialli	Top Hit Descriptor	Homo sapiens EphA4 (EPHA4) mRNA	Homo sapiens v-raf murine sarcoma viral oncogene homolog B1 (BRAF) mRNA	AU117241 HEMBA1 Homo saciens cDNA clone HEMBA1000968 5	Homo sapiens RIBIIR gene (partial), exon 12	Homo saplens actin-related protein 3-beta (ARP3BETA), mRNA	qe62h01.x1 Soares_fetal_bung_NbHL19W Homo sapiens cDNA clone IMAGE:1743601.3' similar to gb:L11566 60S RIBOSOMAL PROTEIN L18 (HUMAN):	qe92h01.x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1743601 3' similar to db:L11566 60S RIBOSONAL PROTEIN 18 (HIMAN)	od60a03.s1 NCI OGAP GCB1 Homo saniens cituA clone IMA GE-1372300 3'	wm26h11.x1 NCI CGAP Ut4 Homo saniens cDNA clone IMAGE-2437195.3	601344705F1 NIH MGC 8 Homo saplens cDNA clone IMAGE:3677641 5	wh57b06.x1 NCI_CGAP_Kld11 Homo sepiens cDNA clone IMAGE:2384819 3' sImilar to TR:055137 055137 ACYL-COA THI0ESTERASE.	Homo saplens latent transforming growth factor beta binding protein 2 (LTRP2) mRNA	Homo sapiens latent transforming growth factor beta binding protein 2 (1 TBP2) mRNA	AU119634 HEMBA1 Homo sapiens cDNA clone HEMBA1006283 5'	601110371F1 NIH MGC 16 Hamo sapiens cDNA clane IMAGE:3351352 5	Homo saplens Smad- and Olf-interacting zinc finger protein mRNA, partial cds	yd08a02.r1 Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:24880 5' similar to SP:A48836 A48836 SPEGFIII≽EGFREPFAT-CONTAIMING FIRROPELLINJI IKE PROTEIN - SEA I IB OLINI	Homo saplens aconitase 2. mitochondrial (ACO2), mRNA	Homo sapiens dNT-2 gene (or mitochondrial 5/3)-denxyribanucle-ritidasa (dNT-2 gene) owne 1-5	Homo saplens sparm surface protein (HSS), mRNA	Homo sepiens short chain L-3-hydroxyacyl-CoA dehydrogenase precursor (HADHSC) gene, nuclear gene encoding milochondrial partein, complete cats	Homo saplens arm-repeat protein NPRAP/neurolungin (CTNND2) mRNA partial cds	Homo sapiens TRAF6-binding protein T6BP mRNA, complete cds	UI-H-BI1-acw-g-01-0-UI.s1 NCI_CGAP_Sub3 Hamo sapiens cDNA clane IMAGE:2715840.3'	EST88807 HSC172 cells II Home sapiens cDNA 5' end similar to similar to ribosomal protein S18	H.seplens mRNA for N-ecety/glucosamide-(beta 1-4)-galactosy/transferase
מום דייסיין סופ	Top Hit Database Source	L	N FN	EST HUMAN	NT	L	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	LX.	LZ	EST_HUMAN		LZ	EST HUMAN						LN	EST_HUMAN	EST_HUMAN	LN
5	Top Hit Acession No.	4758279 NT	4757867 NT	AU117241.1	AJ237744.1	9966912 NT	Al192764.1	Al192764.1	AA826039.1	AI873630.1	BE561063.1	AI764973.1	57732	4557732 NT	AU118634.1	BE258012.1	AF221712.1	T80514.1	1418185	AJ277557.1	11426786 NT	AF095703.1	U52351.1	AF268075.1	AW138646.1	AA376399.1	X13223.1
	Most Similar (Top) Hit BLAST E Value	9.0E-69	9.0E-69	9.0E-69	8.0E-69	7.0E-69	6.0E-69			4.0E-69/	4.0E-69	4.0E-89	4.0E-69	4.0E-69	4.0E-69	3.0E-69	3.0E-69 ⊿	3.0E-69 T	3.0E-69	3.0E-69 A	3.0E-69	3.0E-69.A		3.0E-89	3.0E-69		3.0E-69 X
	Expression Signal	1.15	0.71	6.6	1.09	6.49	8.09	80'8	1.01	1.76	1.42	5.28	2.71	2.71	0.62	3.69	2.32	3.19	3.64	0.67	0.74	0.68	1.83	8.32	0.88	0.65	1.01
	ORF SEQ ID NO:	27741		L		32008	33581	33582			31384	31467	32301	32302	34651	25848	26008		37797		30563	33032	33082	33219	34106		35152
	Exan SEQ ID NO:		16852	23488	16140	19033	20458		21564	13291	25073	18545	19297	19297	21504	13202	13376	14294	17894	19620	17928	19956	20003	20127	20964	21358	21880
	Probe SEQ ID NO:	2275	4109	10805	3381	6259	7762	7762	8873	507	5674	5753	6531	6531	8812	377	598	845	5163	6705	6759	7272	7320	7451	8270	9998	8313

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		- ≥	5	J	1		1	1	1	1	1	1		Т	Т	Т	Т	Т	Т	Т	Г	Г	Г		Īαί	1	ò	т	_	7	, , ,
Oligie Exult Propes Expressed in Brain	Top Hit Descriptor	Human mRNA for calcium-binding protein in macrophages (MRP-14) macrophage migration inhibition factor	(Mir)-related protein	Indmo sapiens SEC10 (S. cerevisiae)-like 1 (SEC10L1), mRNA	France Sapiens ribosomal protein S15a (RPS15A), mRNA	Homo sources UCCS cells II Homo sapiens cDNA 5' end similar to similar to ribosomal protein S18	Home contract VIA Acres	Home saplens KIAA0553 protein gene, complete cds, and alphalib protein gene, partial cds	Home septiens KIAAU533 protein gene, complete eds, and alphalib protein gene, partial eds	Home control of A A person	Found September N/AAUSS3 protein gene, complete cds; and alphalib protein gene, partial cds	2010011 Street 1811	Zm20401 rt Stratege ustis_NH i Homo sapiens cDNA clone IMAGE:781682 5	Data Consider the Paragraph (#93/208) Homo saplens cDNA clone IMAGE:527088 5	And SA 25 A THE ACCURACY CONTROLL SINGLE CONTROLL CONTROLL DE CONT	RO187578984 I NIT _MCC_21 Homo saplens cDNA clone IMAGE:3635781 5	SO1477505054 AIII- 1100 Septens cDNA clone IMAGE:3958532 5	OVO TTOMO ASSESS ST. From Sapiens cDNA clone IMAGE:3958532 5	Home series (1/A 6246	Homo sapiens KIAA0718	Homo septens mRNA for KIAA4442	Homo saplens mRNA for KIAA147 motern, partial cds	601278532F1 NIH M.C. 30 Home cont. Partial cds	601278532F1 NIH MGC 38 Homo sapilans cDNA clone INA GE:3810814 5	TCBAP1E2678 Pediatric pre-B cell scute lymphoblastic leukemia Baylor-HGSC project≕TCBA Homo saplens cDNA rione TCBA page 7	TCBAP1E2978 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC projecte TCBA Homo contract	cDNA clone I CBAP2678	Homo sapiens mRNA for KIAA0707 protein, partial cds	602043782F1 NCI_CGAP_Bm67 Home sapiens cDNA clone IMAGE:4181325 51	Homo saplens keratin 8 (KRT8) mRNA	WITHOUSE SOURS NIT. T GBC, 20 Home saplens cDNA clone IMAGE:4025785 6' WITHOUSE SOURCE NIT. T GBC, S1 Home saplens cDNA clone IMAGE:2360360 3' similar to contains Alu
ואום ביאחוו גדונ	Top Hit Database Source	17	1	Z L	ECT DIMAN	-1	Į	Į.	Į	Į.	EST HUMAN	FST HIMAN	EST HUMAN	Į.	EST HIMAN	EST HIMAN	EST HIMAN	FST HIMAN	-1	: -	N-	LN	EST HUMAN	EST_HUMAN	EST HUMAN	T	NAMOL	Ī	HOMAN	T. CLINARY.	T
5	Top Hit Acession No.	-69 X08233 1	F720000		AA376300 1	3.0E-69 11419157 NT	AF1602521	2.0E-69 AF160252 1	Τ	Γ				Γ	T	Γ	Γ	1.0E-69 AW393969.1	2283	7662263 NT	69 AB032973.1	89 AB032973.1 IN	59 BE531007.1 E	39 BE531007.1 E	1.0E-69 BE245070.1		Ī		200	2	
	Most Similar (Top) Hit BLAST E Value	3.05	308	306			١	ı			2.0E-69	2.0E-69	2.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-89	1.0E-69/	1.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-69	1.0E-69 B	1.0E-69 B	1 0F-89 B	4 05 80 4	100 00 0	1.0E-69 Br 320429.1	1.0E-69 RF125887 4	1.0E-69 Al809994.1
	Expression Signal	2.03			7.81	5.17	1.64	1.64	10.33	10.33	2.08	3.16	1.08	1.98	0.74	0.87	0.87	4.37	1,26	1.26	3.01	3.01	1.14	1.14	4.91	4.91	1 30	0 67	10.62	1.74	4.45
	ORF SEQ ID NO:	35286	L	36468		Ц					27327		34284	27139		31700	31701	32274	32679	32680	32603	32604	32554	32555	35937	35938	36034	38177	-	37281	
L	SEQ IO	22111	22380								- 1	- 1	- 1	- 1	17887	18741	18741	19273	19836	19636	19571	19571	19526	19528	22721	22721	22818	22961	23465	24512	24769
	Probe SEQ ID NO:	9433	9729	10538	10745	12024	128	126	395	382	1878	2848	8452	1698	4962	5959	2929	8208	6721	6721	6737	18/3	0782	78/0	10073	10073	10168	10314	10782	11964	12366

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_			_	_	_	_	_	_		_	_				_		_		_					_	_						
	Top Hit Descriptor	nc13d12,r1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1008023	Homo sapiens DGS-I mRNA, 3' end	tm89f01.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2165305 31	tm89f01.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2165305 3'	z15h04.r1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:713239 5	Homo saplens tumor suppressor deleted in oral cancer-related 1 (DOC-1R) mRNA	Homo sapiens adenylate cyclase 3 (ADCY3) mRNA	Homo sapiens MIST mRNA, partial cds	Homo sapiens MIST mRNA, partial cds	Homo sapiens gene encoding splicing factor SF1, exons 2-8	Homo sapiens mRNA for KIAA1284 protein, partial cds	Homo sapiens mRNA for KIAA1294 protein, partial cds	Human displacement protein (CCAAT) mRNA	Human displacement protein (CCAAT) mRNA	Human PBX3 mRNA	Human PBX3 mRNA	Homo sapiens phospholipid scramblase 1 gene, expn 1 and 5' flanking region	Homo sapiens karyopherin beta 2b, transportin (TRN2), mRNA	Homo saplens karyopherin beta 2b, transportin (TRN2), mRNA	Homo sapiens glutamate-cysteine ligase (pamma-glutamylcysteine synthetase), catalytic (72.8kD) (GLCLC) mRNA	Homo sapiens NDST4 mRNA for N-deacet/lase/N-sulfotransferase 4, complete cds	Homo sapiens NDST4 mRNA for N-deacetylase/N-sulfotransferase 4, complete cds	Homo saplens spastic paraplegia 4 (autosomal dominant; spastin) (SPG4), mRNA	Homo saplens spastic paraplegia 4 (autosomal dominant, spastin) (SPG4), mRNA	Homo saplens HIR (histone cell cycle regulation defective, S. cerevistae) homolog A (HIRA), mRNA	Homo sapiens HIR (histone cell cycle regulation defective, S. cerevislae) homolog A (HIRA), mRNA	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-tl, Alzheimer disease) (APP), mRNA	Human Ku (p70/p80) subunit mRNA, complete cds	Homo saplens CMP-N-acetylneuraminic acid synthase (LOC55907), mRNA	Homo sapiens KIAA0792 gene product (KIAA0792), mRNA
<u></u>		EST_HUMAN	N N	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝŢ	IN	N	N	N	NT	N	F	۲N	ΡN	LN	NT	F E	Į.	Ä	LN.	۲	NT	NT	TN	TN	LN	١	N L	۲N
	Top Hit Acession No.	70 AA230303.1	70 L77566.1	70 AI497807.1	70 AI497807.1	70 AA282955.1	5031668 NT	4767723 NT	70 AB032369.1	70 AB032369.1	70 AJ000052.1	70/AB037715.1	70 AB037715.1	70 M74099.1	70 M74099.1	70 X59841.1	70 X59841.1	70 AF153715.1	11525964 NT	11525964 NT	4557624 NT	70 AB036429.1	7.0E-70 AB036429.1	11429685 NT	11429685 NT	11526319 NT	11526319 NT	4502166 NT	M30938.1	TN 8923899 NT	7662307 NT
	Most Similar (Top) Hit BLAST E Value	8.0E-70	8.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	7.0E-70	6.0E-70	6.0E-70 M30938.1	6.0E-70	5.0E-70
	Expression Signal	2.08	1.93	1.01	1.01	1.66	2.92	3.67	4.88	4.88	2.16	2.2	2.2	3.58	3.58	3.8	3.8	3.43	2.69	2.69	0.57	0.62	0.62	1.54	1.54	1.65	1,65	1.77	1.21	1.22	2.18
	ORF SEQ ID NO:	27791						29566		30889	32509					34902			133551	33552	35394		36059		36942	37515	37516	26291	27593		28003
	Exon SEQ ID NO:	15591	17079	14546	14546	14660	14788	i	18195	18195	19487	21023		21314	21314			20406		20431	22209	1	22844	.	23682	24196	24196	13621	14863		15598
	Probe SEQ ID NO:	2331	4340	1806	1806	1923	2056	4189	5395	5395	6826	0658	8330	8622	8622	9055	9022	9335	9361	19861	9296	10196	10196	11010	11010	11597	11597	851	2133	2513	2555

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		Homo sapiens KIAA0792 gene product (KIAA0792), mRNA	MR3-H10487-150200-115-a06 HT0487 Hamo sapiens cDNA	IES I 03928 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBDN26	ICM4-UM0003-010300-105-g08 UM0003 Homo sapiens cDNA	CM4-UM0003-010300-105-g08 UM0003 Homo sapiens cDNA	RCU-B 10522-071299-011-a12 BT0522 Homo saplens cDNA	r.c5 l 0222-011299-011-a12 BT0522 Homo sapiens cDNA	Homo sapiens plakophilin 4 (PKP4), mRNA	Homo seplens plakophilin 4 (PKP4), mRNA	whey doubt a NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:2388005 3'	602741561F1 NIH_MGC_46 Homo saplens cDNA clone IMAGE:4302808 5'	602141661F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4302806 5'	hz81h02.x1 NCL_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3214419 31	Homo saplens phosphatidylinositol 4-kinase 230 (ni4k23a) mBNA	yy07a10,r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:270622 5' similer to	VY DOTAL TALL TALE TO 3-HYDROXYISOBUTYRATE DEHYDROGENASE PRECURSOR; VYO7a10.r1 Soares melanocyte 2NbHM Homo saplens CDN4 close MA CE-272/2020	SW:D3HI_RAT P29266 3-HYDROXYISOBUTYRATE DEHYDROGENASE PRECLIDED.	qx51h01.x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2004913.31	Fromo saplens hypothetical protein FLJ20768 (FLJ20768), mRNA	Fromo sapiens KIAA0193 gene product (KIAA0193), mRNA	From Sapiens KIAA0193 gene product (KIAA0193), mRNA	2/48g04.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:380214.6; climitar to CMXCAO LIZZE	P03345 GAG POLYPROTEIN;	ypospu4.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:191599 5/	Human nonmuscle myosin heavy chain-B (MYH10) mRNA, partial cds	n septens gene for schwannomin (CS8)	H.Sapiens gene for schwannomin (CS8)	Fromo saptens NALP1 mRNA, complete cds	Home services and the protein Isoform (neurafibromin Isoform), complete cds	Homo saplens cydnasmic dynein infermediate chain 1 mRNA, complete cds	Sylphasilia dylicin intermediate chain 1 mRNA complete cde
gle Exon Pro	Top Hit Database Source		. 1	EST_HUMAN	EST HUMAN	EST HUMAIN	ESI HUMAN	EST LINAN	HI TOWN	1	11.000	Т	T	Т	HOMAN	LN L	EST HIMAN	7-	7	9000000 LEST HUMAN	Т			T	ES HOMAN	LICIMEN							
Sin	Top Hit Acession No.		5.0E-70 FE46600 / 662307 NT	T		T	T	T	0000	1143099911	70 AIR31074 4		T	T	70 4 5040070 4		70 N42161.1			S S S S S S S S S S S S S S S S S S S	7664083 NIT	7661083 NIT	70 AL 163202 2 NT		T			T	-	T	Ţ.		
	Most Similar (Top) Hit BLAST E Value	20 20	0.0E-70	4 0F-70	4.0E-70	4.0F-70	3.0E-70	3.0E-70	3.0E-70	3.0E-70	3.0E-70	3.05-70	3.0E-70	3.0F-70	205	Z.UE-/U/	2.0E-70	1000	2 05 70 4	2.0E-70	20E-70	2.0E-70	2.0E-70 A	2 0F.70 AA054040 4	20E-70H	2.0E-70 M69181 1	2.0E-70 X	2.0F-70 X	2.0E-70 AF	2.0E-70 D	2.0E-70 AF	2.0E-70 AF	
	Expression Signal	2 18	2 Z	1.03	1.78	1.78	1.23	1.23	0.65	0.65	1.6	1.65	1.65	0.58	12	!	14.09	4 00	1,12	1.56	3.05	3.05	1.66	6.32	0.78	5.06	8.7	8.7	1.27	1.75	12.14	12.14	
	ORF SEQ ID NO:	28004		32431	32663	32654	27016	27017	31234	31235	31581	32031	32032	35869	25484		26089	26090	26113	26421	26578	26579	27177		29011	29404	30839	30940	31857	32307	32342	32343	
	Exon SEQ ID NO:	16598	L	19417	H	19613	14330				18642			22656	12865		13449	13449	13464	13760	13915	13915	14478	15043	16368	16772	18227	18227	18888	19303	19333	19333	
	Probe SEQ ID NO:	2555	11974	9999	9699	9699	1584	1584	5532	5532	5855	6280	6280	10008	37		673	673	88	1000	1161	1161	1736	2318	3615	4027	5428	5428	6111	6538	6269	6269	
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Table 4
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	. Top Hit Descriptor
6897	17973		1.68	2.0E-70	11422642 NT	_ LN	Homo saplens sialytransferase 6 (N-acetyllacosaminide alpha 2,3-stalytransferase) (SIAT6), mRNA
7303	19986	33062	97.0	2.0E-70	70 AF288207.1	١	Homo sapiens cysteiny-tRNA synthetase mRNA, complete cds, alternatively spliced
7819	20514	33639	9.02	2.0E-70	70 M21741.1	NT	Human guanine nucleotide-binding protein alpha-subunit gene (G-s-alpha), exons 4 and 5
8121	20815	33951	0.5		11423599 NT	TN	Homo sapiens amyto-1,6-glucosidase, 4-alpha-glucanotransferase (glycogen debranching enzyme, glycogen storage disease type iii) (AGL), mRNA
8228	21250		0.8	2.0E-	70 H47959.1	EST_HUMAN	yp79g02.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:193682 5'
9067	21756		0.95	2.0E-70	11526355 NT	LN T	Homo sapiens dynactin p62 subunit (LOC51164), mRNA
10038	22686		1.46	2.0E-70		NT	Homo sapiens calcium-binding transporter mRNA, partial cds
10500	23146	36373	0.47	2.0E-	70 AB033042.1	ΤN	Homo sapiens mRNA for KIAA1216 protein, partial cds
11005	23677	36933	3.75	2.0E-7	8923420 NT	NT	Homo sapiens hypothetical protein FLJ20450 (FLJ20450), mRNA
11005	23677	36934	3.75		8923420 NT	ΙN	Homo sapiens hypothetical protein FLJ20450 (FLJ20450), mRNA
11639	24236	37559	7.32		4503520 NT	NT	Homo saplens eukaryotic translation initiation factor 3, subunit 6 (48kD) (EIF3S6) mRNA
12353	24757	31058	3.06		11430460 NT	LΝ	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
12353	24757	31059	3.06	2.0E-70	11430460 NT	ΙN	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA
3388	16147		2.63	1.0E-70	TN 974767	FZ	Homo saptens transglutaminase 3 (E polypeptide, protein-glutamine-gamma-glutamyftransferase) (TGM3) mRNA
9180	21850		0.69	1.0E-70	70 W85795.1	EST HUMAN	zh55g05.r1 Soares fetal liver spleen 1NFLS S1 Homo sapiens cDNA clone IMAGE:416024 5
8698	22349		0.65	1.0E-70	-	EST HUMAN	zv54c03.r1 Soares testis, NHT Homo sapiens cDNA clone IMAGE:757444 5
10852	23532	36777	7.13	1.0E-70	Γ	EST_HUMAN	AV738538 CB Homo sapiens cDNA clone CBLBGB10 5'
5854	18641	31579	7.05	9.0E-71	9.0E-71 AI143870.1	EST HUMAN	qe04f01x1 Soares testis, NHT Homo sapiens cDNA clone IMAGE:1738009 3' similar to TR:014045 O14045 PHOSPHOTRANSFERASE.
5854	18641	31580	7.05	9.0E-71	9.0E-71 AI143870.1	EST HUMAN	qe04f01x1 Soares, tastis, NHT Homo sapiens cDNA clone IMAGE:1738009 3' similar to TR:O14045 O14045 PHOSPHOTRANSFERASE;
6932	19668	32714	2.23	9.0E-71	1 AI654903.1	EST_HUMAN	wb52c05x1 NCI_CGAP_GC8 Homo saplens cDNA clone IMAGE:2309288 3' similar to TR:P97213 P97213 CDU2, CDU1, TCDD, TCDB, TCDE, TCDE, TCDC, CDD1, CDD2, CDD3, AND CDD4 GENES.
11508	19668	32714	4.79	9.0E-71	9.0E-71 AI654903.1	EST_HUMAN	wb52c05x1 NCI_CGAP_GC8 Home saptens cDNA clone IMAGE:2309288 3' similar to TR:P97213 P97213 CDU2, CDU1, TCDD, TCDB, TCDE, TCDE, TCDC, CDD1, CDD2, CDD3, AND CDD4 GENES.
8968	21658		5.03	8.0E-71	8.0E-71 AA171451.1	EST HUMAN	zp21d11.r1 Stratagene neurospithelium (#837231) Homo sapiens cDNA clone IMAGE:610101 5' similar to TR:01143061 G1143061 STRAIN XA34 POL:
7275		33036	8.9	7.0E-71		Γ	zv60h06.r1 Soares_testis_NHT Hamo sapiens cDNA clone IMAGE:758075 5'
8578	l	34408	1.02	7.0E-71		EST_HUMAN	291a06.s1 Scares_fetai_liver_spleen_1NFLS_S1 Homo sepiens cDNA clone IMAGE:462226 3'
11302	ı	37262	2.07	7.0E-71			Homo sepiens chromosome 21 segment HS21C010
2207	14935	27673	5.97	5.0E-71		LN	Homo sepiens SP100-HMG nuclear autoantigen (SP100) mRNA, complete cds

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		_	Т	Т	Т	T-	T	т-	Τ-	_	_	_	_	-	, -		_			_	_	_	_										
Single Exon Probes Expressed in Brain	Top Hit Descriptor	· · · · · · · · · · · · · · · · · · ·	QV4-51 0234-181199-037-t05 ST0234 Homo sapiens cDNA	Indiana saplens cyclim-dependent kinase 6 (CDK6) mRNA	nomo sapiens keratin, hair, acidic, 7 (KRTHA7), mRNA	Home sapiens MA40623 gene product (KIAA0623), mRNA	From September 101 NAVA1280 protein, partial cds	Home carlons and clarifications of the contraction	Human paringhymathala anthia 4	Homo sariene MA DI IV section 1996 T I TRINA, 3 end of cds	Home seniors become for promise 1991; From Associated with Lins 2 (LOC51678), mRNA	Homo earliers utanscription ractor WS I F mRNA, complete cds	Home septens nuclear ractor related to kappa B binding protein (NFRKB) mRNA	Unition Brook a more at record related to kappa B binding protein (NFRKB) mRNA	Homo sanlens ICE II mDN/k http://www.com/cells	Homo septems non-valuable hosts and the contract of the contra	tissue-activating peofide III, neutronhil-activating pages of 2000 p.	Homo saplens similar to hymothetical probable [120422] (Fr DF), mKNA	Equits caballitis attracted above to the control of the appeals (LOC63325), mRNA	Eduta caballita olyneraldahuda 2 phonta da da da da da da da da da da da da da	Homo sapiens plasminoses (DLC) DNA	Homo saplens SP400-HMG nuclear or thought and the saplens SP400-HMG nuclear or thought and the saplens space a	Homo saplens putative heme-binding protein (SOI II)	down canian actual to the control of	All 1425224 DI ACE 11.	ni45h10.s1 NCI_CGAP_Pr4 Homo sapiens cDNA clone IMAGE:1043883 similar to contain ETDE to ETDE.	repetitive element;	Homo sapiens chromosome 21 segment HS21Cnns	Human mRNA for KIAA0222 gene nertial cde	Human mRNA for KIAAA772 cano consideration	DKFZb434D1721 11 434 (swamm: https://lim	7785511 x1 NOL CGAP_OVID Him subsets that septems cDNA done DKFZp434D1721 6' 7785511 x1 NOL CGAP_OVID Home septems cDNA done IMAGE:3571221 3' similar to TR:Q92165 092165 PUTATIVE FOUR DEDEAT FOR CAMPINAL CONTRACTORS	CONTRACTOR SELECTION CHANNEL:
igle Exon Pro	Top Hit Database Source	FOT LI MANN	NAMOR INCIDENT	Z E	Z E	i k	Į.	LN.	LN	L	LN	Ę	Į.	LN	TA L		N	LZ	IN.	NT	5	¥	5		T HUMAN	\top	T_HUMAN			LN LN	T HUMAN		7
	Top Hit Acession No.	71 AW818405 1	4502740 NIT		TAROCCART	5.0E-71 AB033108.1	AB033108 1	11431590 NT		11526445	AF072810.1	377	5453777 NT	5.0E-71 X13487 1	5729900		11436514 NT	11438069 NT		-71 AF157626.1	5880	-71 AF056322.1	7602	7019352 NT	71 AU135734 1			3.2	71 D87462.1	71 D87462.1	-		
	Most Similar (Top) Hit BLAST E Value	5.0E-71	5.0F-71	5.0E-71	5 0F-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71	5.0E-71		5.0E-71	5.0E-71	4.0E-71	4.0E-71	4.0E-71	4.0E-71	4.0E-71	4.0E-71	3.0E-71	100		2.0E-71 A	2.0E-71 D	2.0E-71 D	20E-71 A	20E-71B	
	Expression Signal	1.36	2.23	1.42	1.43	0.62	0.62	0.76	1.47	0.75	22.56	0.61	0.61	2.67	1.45		2.93	2.57	102.7	102.7	1.97	3.37	6.57		1.41		300.50	2.02	8.24	8.24	0.55	0.64	
	ORF SEQ ID NO:	28472	L	L	Ĺ	LI					33408	34261	34252		38461		36831	37097	25778	25779	28299	29762	30288	30404		36503	2000	25626	30672	30673	30543	34735	
	SEQ ID	16844	18581			l		j	IJ	J	ı	21114	21114	22462	23227		23581	23814	13141	13141	15658	17131	17684	17788	20628	23285		2000	200	18043	17948	21694	
L	Probe SEQ ID NO:	4101	5790	6564	6821	6878	6878	7046	7410	7607	788	8421	8421	9811	10530			7	좕	8 8 8	7888	4384	4926	6909	7933	10591	138	2007	1000	923/	6871	8903	

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		Homo sapiens glypican-6 (GPC6) mRNA, complete cds	Homo sepiens myomesin (M-protein) 2 (166kD) (MYOM2), mRNA	Homo sapiens hypothetical protein FLJ10998 (FLJ10998), mRNA	Fromo sapiens hypothetical protein FLJ10698 (FLJ10998), mRNA	CSNKZA1≖casein kinase II (CKII) subunit alpha [human, Genomic, 18862 nt]	nomo sapiens cytochrome c oxidase subunit VIIa-related protein gene, complete cds	AVOIST WILS HOME SEPIENS CDNA clone MDSE(A03 6)	AV784347 N.C. I.	AV701217 MIDS Home sapiens cDNA clane MDSEIA03 6'	From Septens coagulation factor XIII, A1 polypeptide (F13A1), mRNA	normo saprens reucy//cystiny/ aminopeptidase (LNPEP), mRNA	nomo sapiens leucyl/cystinyl aminopeptidase (LNPEP), mRNA	inomo saptens gene for AF-6, complete cds wk95g03.x1 NCI_CGAP_Lu19 Home saptens cDNA_close IMACE:2422466.21	HYPOTHETICAL 38.6 KD PROTEIN, contains Alu repetitive element:	wk95g03.x1 NCI_CGAP_Lu19 Homo sapiens cDNA clone IMAGE:2423188 3' similar to TR:088705 088705	ANTICINETICAL 38.6 KD PROTEIN. ;contains Alu repetitive element;	ou 1456/4/71 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3862451 6	Homo saplens aconitase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial protein, mRNA	Homo saplens aconitase 2. mitochondrial (ACO3), modoz	M. C.C., morest gene encoding milocondrial protein, mRNA	numo sapiens aconitase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial protein mRNA	pseudogene) PTMAP2=prothymosin alpha [human, Genomic, 1192 nt. segment 2 of عالم المرابعة ال	HSPD13670 HM3 Home saplens cDNA clone s4000061G02	Homo sapiens chromosome 21 segment HS210046	QV0-CS0010-150900-398-e11 CS0010 Homo seplens cDNA	QV0-CS0010-150900-398-e11 CS0010 Homo sapiens cDNA	QV0-CS0010-150900-398-e11 CS0010 Homo saplens cDNA	QV0-CS0010-160900-398-e11 CS0010 Hamo saplens cDNA	Homo saplens alpha-tubulin mRNA, complete cds	AU128584 NT2RP2 Homo saplens cDNA clone NT2RP2003761 6'
jle Exon Probe	Top Hit Database Source							T HI MAAN	NONCH I	T LI BAANI	NAME					EST_HUMAN HY	EST HIMAN LIVE	T	NEW					7	T_HUMAN					T_HUMAN	F	EST_HUMAN AU1
Sin	Top Hit Acession No.	74 05405007 4	18	11425430 N	09/2/811 N I	1 572303 4	-		3142	71 AV761217 1	Ronz	41447494INT	T1417101 INT	1 AB011300 1	T	2 Al857635.1 E	9.0E-72 AI857635 1 F			4501866 NT	4501866 NT	4E01BER	C446044							-		AU128584.1 ES
	Most Stmitar (Top) Hit BLAST E Value	1 OF 74	4 OF 74	1 0E-74	1.0F-74				1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71	1.0E-71		9.0E-72	9.0E-72	8.0E-72		7.0E-72	7.0E-72	7.0E.72	7 OF 79 C		7.05-721		5.0E-72B	6.0E-72B	6.0E-72 B	9.0E-72.B	5.0E-72 L	D.UE-72IA
	Expression Signal	0 6	2.26	4 09	408	0.78	7.89	2.05	1.45	2.58	2.19	1.73	1.73	4.96		1.72	1.72	0.97	8	6.9	2.63	2.63	280	100	01 /	1 6	90:-	1.06	3.47	0.47	3.62	80.
	ORF SEQ ID NO:	33872	33891			34982	35752		36309		36720	37037	37038		00000	73830	25831	31762	20.450	2	29459	29460	32773		-	26824	25527	25524	25522	2002	225B4	10000
	Exon SEQ ID NO:	20740	ı				22557		23082	23375	23479	23763	23763	24786	12183	3	13183	18801	16834		16834	16834	19716	24868	20977	12880	12880	12880	12880	13874	19551	
	Probe SEQ ID NO:	8046	8069	8345	8345	9128	8066	8968	10438	10684	10796	11093	11093	12401	308	8	398	6020	4092		4082	4092	7024	12520	8283	90	e G	3 6	8	1117	6851	

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Table 4
Single Exon Probes Expressed in Brain

Top Hit Descriptor	EST188312 HCC cell line (matastasis to liver in mouse) Il Homo sapiens cDNA 5' end similar to similar to FAC1	au80c03.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782564 5' similar to TR:099785 099785 HYPOTHETICAL 32.4 KD PROTEIN ;contains element MSR1 repetitive element ;	Г	MR4-BT0598-010600-005-d05 BT0598 Homo sapiens cDNA	NR4-BT0598-010600-005-d05 BT0598 Homo sapiens cDNA	V ba08g08.y1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2823806 5'		Г	Г		Homo saplens heet domain and RLD 2 (HERC2), mRNA	Homo sapiens hypothetical protein FLJ20758 (FLJ20758), mRNA	V RC3-LT0023-200100-012-d11 LT0023 Homo sapiens cDNA	V RC3-LT0023-200100-012-d11 LT0023 Homo saplens cDNA	qh67c02.x1 Soares, fetal liver, spleen_1NFLS_S1 Homo saplens cDNA clone IMAGE:1849730 3' similar to TR:014498 Q14498 SPLICING FACTOR. [1] contains Alu repetitive element contains element L1 repetitive		lea23f09.s1 NC_CGAP_GCB1 Homo saplens cDNA clone IMAGE:814121 3' similar to SW:CPTR_FLAPR P9131 CHLOROPLAST TRIOSE PHOSPHATE TRANSLOCATOR PRECURSOR.;	aa23f09.s1 NCI_CGAP_GCB1 Homo septens oDNA clone IMAGE:814121 3' similar to SW:CPTR_FLAPR	Т	Т	Homo sapiens eukaryotic translation Initiation factor 2B, subunit 2 (beta, 39kD) (EIF2B2), mRNA	yd28d09.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:1096493'	Homo sapiens WEE1 gene for protein kinase and partial ZNF143 gene for zinc finger transcription factor	Hamo sapiens pre-B-cell colony-enhancing factor (PBEF) mRNA	П
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	F	EST HUMAN	Ę	LN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT	뉟	EST_HUMAN	LΝ	F	EST_HUMAN
Top Hit Acession No.	72 AA316632.1	72 AW161274.1	72 AV724632.1	72 BF331571.1	72 BF331571.1	72 BE208545.1	72 BE208545.1	72 BE926645.1	72 AF170025.1	72 T87947.1	5729867 NT	8923669 NT	72 AW836230.1	72 AW836230.1		72 AI248796.1	72 AA465388.1	72 AA465388.1	72 H79421.1	7657057 NT	7657057 NT	72 T81910.1		3.0E-72 5031976 NT	AA723823.1
Most Similar (Top) Hit BLAST E Value	5.0E-72	5.0E-72	5.0E-72	5.0E-72	5.0E-72	5.0E-72		6.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72		4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	4.0E-72	3.0E-72	3.0E-72
Expression Signal	0.73	3.71	0.89	3.45	3.45	1.61	1.61	2.82	0.62	89:0	2.03	1.3	0.48	0.48		0.92	1.57	1.57	7.78	1.75	1.75	2.16	8.92	69'0	1.52
ORF SEQ ID NO:	33509	34514	35708	37157			37564		30868	32230	33069		36152	36153		36186	37210	37211	L		37556	37597	31025		
Exen SEQ ID NO:	20394	21367	22511		23871			25358	18178	19229	19992	22336		22839		22967	11682	23917	ł	24234	24234	24275	24822	l	1 1
Probe SEQ ID NO:	7731	8675	9861	11208	11208	11643	11643	12107	5378	6462	7309	9684	10292	10292		10320	11255	11255	11514	11637	11637	11680	12453	18	883

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						21 1127 218	Charle Librar Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
1132	13888	28546	6.64	3.0E-72	72 016306.1	칟	Human chondroith suifate protecolytean versitean V/O enlinearminant
1132	13888	26547		3.0E-72	U16306.1	5	Himan characters affice and the control of the cont
1171		26587	0.72	3.0E-72	-72 U80226.1	Ę	Human gamma-aminohirhyir anid fransomhang and fransomhang and the control of the
1171	13925	26588		3.0E-72	72 U80226.1	Ā	Human gamma-aminobutvic acid transaminase mRNA northings
1510	14256	26942	1.24	3.0E-72	BE242181.1	EST HEMAN	TCAAP1E1252 Pediatric acute myelogenous leukemia cell (FAB M1) Baylor-HGSC project=TCAA Homo
3072	15838	28481	11.45	3.0E-72	3.0E-72 AJ229043.1	LN	Homo saniane 950 kh Amilia hallona a Milia - 1 Amilia
3273	16034	28684	2.17	3.0E-72	8923548	Ę	Homo saplens bunchaited broken FI Tobase (FI Tobase)
3803	16555	29186	2.69	3.0E-72	3.0E-72 S77589.1	FN	TCR V delta 2-C alpha = T-cell receptor delta and C alpha fusion gene (alternatively spliced, splice function)
4508	17243	29876	3.12	3.0E-72	1416198	Į	Francist, productor Decement Ren, mrnv Parial, 211 ntj. Homosaniens handhaffing broth 751 144473, 19444
4715	17447	30079	1.07	3.0E-72	3.0E-72 AF167572.1	LZ	Homo sapians profein methyltransference (1954), mithyl
4715	17447	30080	1.07	3.0E-72		LN	Homo sapians protein methyltranderno (1954) - DNA
4862	17591	30215	0.95	3.0E-72		T HUMAN	wb31a08 x1 NCI CG4P GC8 Home continue that the c
5433	18232		1.27	3.0E-72	9083		domo sanjene samanboh W CEMMM - This Color Galle IMAGE:230/254 3
5891	18676	31621	2.1	3.0E-72 /	3.0E-72 AF073367.1		Homo saplens growth factor recently have a series of Anna a
5891	18676	31622	2.1	3.0E-72	Π		John Saplens growth forder country. A contract of CRETO gene, exch 5
6074	18853	31819	4.82	3.0E-72	T		tremin september 3 your unique technicability of GRB10) gene, exon 5
6074	18853	31820	4.82	3.0E-72 A			Homo septementally for It A A 2001
6516	19281	32284	3.63	3.0E-72	2869		Homo septions ribosome in rolein 13 like (RB) 31 mb/l A
7485	20157	33249	2.15		U80017.1	F	Homo sepiens basic transcription factor 2 p44 (bff2p44) gene, partial cds, neuronal apoptosis inhibitory
8075	20769	33898	0.98		5031892		Homo seniens hucher recentor su Memily 4 min U
10331	22978	36198	1.2	3.0E-72 X	X98289.1		Homo sapiens \$100412 pane for Celebraniile Common 2 11:11
5869	18656	31597	1.91	2.0E-72	1426671		Homo sapiens solute carrier family 13 (sodium-dependent dicerboxylate transporter), member 2 (SLC13A2), mRNa.
8983	21683	34832	0.71		BF308560.1	H IMAN	THE WAS AT HE WAS 47 H.
8993	21683	34833	0.71	2.0E-72 B	Ī	Т	601890419F1 NIH MGC 17 Home services CDNA clone IMAGE:4131461 6
10640	23331	36569	2.47	2.0E-72 A	AA789277.1		aj28b09.s1 Soares_testis_NHT Homo septems cDNA clone 1391609 3 similar to gb:X02067 H.sepiens
12449	24819	31022	5.75		T	Т	affile population supplies the contract of the
2068	14800	27527	1.19	1.0E-72 A	T	T HIMAN	eliRed 2 - S. cozze. percenting processing the processing of the p
						7	COULTS! Society Barathyrold Limbor North Homo sapiens cDNA clone IMAGE:1387395 3.

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
5680	18473	31390	3.63	1.0E-72	7657676 NT	NT	Homo sapiens vacuolar protein sorting 41 (yeast homolog) (VPS41), mRNA
6464		32231	1.31	1.0E-72	11321578 NT	ΝΤ	Homo saplens myosin, heavy polypeptide 13, skeletal muscle (MYH13), mRNA
6464	19231	32232	1.31	1.0E-72	11321578 NT	ΝT	Homo saplens myosin, heavy polypeptide 13, skeletal muscle (MYH13), mRNA
6536	1	32305	1.2	1.0E-72	AV751818.1	EST_HUMAN	AV751818 NPD Homo sapiens cDNA clone NPDAIE11 5'
7537		33304	3.7		BE175434.1	EST_HUMAN	RC4-HT0578-170300-012-g02 HT0578 Homo sapiens cDNA
7537		33305	3.7	1.0E-72	2 BE175434.1	EST_HUMAN	RC4-HT0578-170300-012-g02 HT0578 Homo sapiens cDNA
9491	22144	35324	10.25	1.0E-72	2 AF222742.1	NT	Homo sapiens synaptic glycoprotein SC2 (SC2) mRNA, complete cds
9491	22144	35325	10.25	1.0E-72	2 AF222742.1	TN	Hamo sapiens synaptic glycoprotein SC2 (SC2) mRNA, complete cds
1444	14191	26875	1.35	9.0E-73	AW374968.1	EST_HUMAN	MR0-CT0063-071099-002-h11 CT0063 Homo sapiens cDNA
10871	23551		15.11	9.0E-73	11424099 NT	1	Homo sapiens ribosomal protein L13a (RPL13A), mRNA
1015	13774	26434	2.29		AW071755.1	EST HUMAN	ws55c06.x1 NCI_CGAP_Brr25 Homo sapiens cDNA clone IMAGE:2501098 3' similar to TR:Q59050 Q59050 HYPOTHETICAL PROTEIN MJ1656. ;
5493	18292	31190	-	8.0E-73	862	IN	Homo sapiens phosphattdylmosttol 3-kinase, class 2, alpha polypeptide (PIK3C2A) mRNA
6478	19245	32245	5.16	8.0E-73	11426469 NT	NT	Homo sapiens lysozyme homolog (LOC57151), mRNA
7994	50689	33817	2.87	8.0E-73	AF113129.1	N	Homo sapiens vecuolar ATPase isoform VA68 mRNA, complete cds
9253	21932	35105	6.25	8.0E-73	BE019900.1	EST HUMAN	bb62a08.y1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3030034 5' similar to gb:X04098_cds1 ACTIN, CYTOPLASMIC 2 (HUMAN); gb3M21495 Mouse cytoskeletal gamme-actin mRNA. complete cds (MOUSE):
9640	<u>L</u>	35484	1.92	-	26037		Homo sabiens Interleukin 12 receptor, beta 1 (IL12RB1), mRNA
9640	L	35485	1.92	8.0E-73	11526037 NT	IN	Homo saplens Interfeukin 12 receptor, beta 1 (IL12RB1), mRNA
10507	Ш	36379	0.45	8.0E-73	4507628 NT	LN	Homo saplens transition protein 1 (during histone to protamine replacement) (TNP1) mRNA
11573	L_I		1.28	8.0E-73	11418786 NT	TN	Homo sapiens DEAD-box protein (HAGE), mRNA
12506	3 24859		3.31	8.0E-73	11418189 NT	NT	Homo saplens thyroid autoantigen 70kD (Ku antigen) (G22P1), mRNA
1112		26528	68.0	7.0E-73	1N 06282300 NT	NT	Homo sapiens hypothetical protein FLJ20309 (FLJ20309), mRNA
3295	16056	28705	1.18		AL163206.2		Homo sapiens chromosome 21 segment HS21 C006
4891	17618		1.35	7.0E-73	AL163282.2	LN	Homo saplens chromosome 21 segment HS21 C082
154	12969		3.07	6.0E-73	AL163218.2	LN TN	Homo saplens chromosome 21 segment HS21C018
7072	19783	32827	3.48		BE166574.1	EST_HUMAN	QV0-HT0494-020300-137-d03 HT0494 Homo sapiens cDNA
5173		30497	2.2		11422159 NT	TN	Homo sapiens HELG protein (FAM4A1), mRNA
1318	14067	26741	2.77	3.0E-73	AW843789.1	EST_HUMAN	CM0-CN0044-260100-164-f08 CN0044 Homo sapiens cDNA
6598	18361	32374	0.71	3.0E-73	AA136403.1	EST HUMAN	zn95e04.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:565950 3' similar to gb:Z23064, cds1 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN G (HUMAN);
8656	21348	34492	0.65			EST_HUMAN	AV729428 HTC Homo sapiens cDNA clone HTCAAF071 5'
8656	11	34493	0.65	3.0E-73		EST_HUMAN	AV729428 HTC Hamo sapiens cDNA clone HTCAAF071 5'

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	$\overline{}$	_		_	_	_	_							_						_	_			_	_		_					
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Out 1402 XI Scares NE T GBC St Home conjess ADMA along May OF 1608 PM T GBC St Home conjess and along the second statement of	Hama sablens chromosome 24 segment HQ24 OAA	RC3-NN0066-270400-011-c04 NN0066 Homo sanians chiva	Homo sapiens BASS1 (BASS1) mRNA narial cde	RC3-NN0088-270400-011-004 NN0088 Homo saniens cDNA	Human beta globin region on chromosome 11	Homo saplens caspase 8, apoptosis-related cysteine protesse (CASPB) mRNA	Homo saplens Parkinson disease (autosomal recessive, juvenile) 2, parkin (PARK2), transcript variant 3, mRNA	Homo saplens Parkinson disease (autosomal recessive, juvenile) 2, parkin (PARK2), transcript variant 3, mRNA	Home caniane chromosome 31 comes at 1100,000	Mus misculus thorres interesting after Union (Call) and	Mila mileguli a thofrend intercenting citizen (Inner Coll.) DNA	Homo sanlens mRNA for KIAA4ad wrond; model of	Homo sanians Interdentin A recent // ABA BNA	Homo sablens Interfacility 4 recents (11.4D)	Hunan perinheral mailin protein 22 mDNA America 24.	Homo seplens mRNA for KIAA 1329 profess manifely and and	Gallus gallus Dach2 protein (Dach2) mRNA complete cds	Gallus gallus Dach2 protein (Dach2) mRNA, complete crts	Homo saplens glutathione synthetase (GSS) mRNA	Homo saplens supervillin (SVIL), transcript variant 1, mRNA	Homo sapiens supervillin (SVIL), transcript varient 1, mRNA	Homo sapiens galactosyteeramidase (Krabbe disease) (GALC), mRNA	Homo sapiens galactosyceramidase (Krabbe disease) (GALC), mRNA	Homo saplens mRNA for KIAA1059 protein, partial cds	RC3-NN0086-270400-011-c04 NN0086 Hamo saplens CDNA	AU121585 MAMMA1 Homo saplens cDNA clone MAMMA1000490 51	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds	CM1-HT0282-111199-042-h10 HT0282 Homo sablens cDNA	9981b07.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1839637 5' similar to contains element MER22 repetitive element:	801276071F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3617105 6"
gle Exon Pro	Top Hit Database Source	EST HUMAN	Į L	EST HUMAN		EST HUMAN	Į.	FZ	Ŋ		Į	LZ	LZ	L	L	L	Ę	FZ	NT	Į,	Ę	나				NT	EST_HUMAN	EST_HUMAN	Г	EST_HUMAN	EST HUMAN	П
Sin	Tap Hit Acession No.	3.0E-73 AI004040.1	3.0E-73 AL 163246.2	-73 AW898081.1	2.0E-73 AF139897.1	-73 AW898081.1	-73 U01317.1	4502582 NT	7669539 NT	7869539	1 163283 2	-73 AF086824.1	Γ		31471	11431471 NT	73 M94048.1	5	Γ	73 AF198349.1	4504168 NT	11496980 NT	11496980 NT	4557812 NT	4557612 NT		73 AW898081.1	73 AU121585.1 E	73 AF198349.1	3 BE151283.1 E		
	Most Similar (Top) Hit BLAST E Value	3.0E-73	3.0E-73	3.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73 N	2.0E-73 A	2.0E-73 A	2.0E-73 A	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73 A	2.0E-73 A	1.0E-73 A	1.0E-73 A	1.0E-73 B	1.0E-73 AI147427.1	1.0E-73 BI
	Expression Signal	1.51	1.5	1.54	1.43	1.78	1.3	3.99	0.91	16.0	18	0.89	0.89	6.27	1.27	1.27	0.68	0.73	0.62	0.52	1.21	1.31	1.31	3.37	3.37	1.82	1.83	1.71	1.12	1.07	1.37	2.95
	ORF SEQ ID NO:				28271			28590	28943	28944	-	32102	32103	32169	32376	32377	33476	33478	35284	35285	36189	36257	36258	36917	36918	36955		27221	27946	32015	35230	36426
	Exon SEQ ID NO:							15940	16294		17138	19114	19114	19158	18363	19383						23040	23040	23062	23862	23692	14674	14618	16205	19039	22061	23195
	Probe SEQ ID NO:	11611	12734	12738	831	1939	2288	3177	3538	3538	4401	6344	6344	6389	9800	9800	7699	7701	9432	9432	10322	1884	1030	/8801	10887	11020	12283	1778	2488	9599	9338	11428

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Single Exon Probes Expressed in Brain

Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
13497	26150	1.06	8.0E-74	4557426	NT	Homo sapiens CD39-like 4 (CD39L4) mRNA
18613	31544	2.2	8.0E-74		TN	Ca2+/calmodulin-dependent protein kinase IV kinase Isoform [rats, brain, mRNA, 3429 nt]
18613	31545	2.2	8.0E-74		NT	Ca2+/calmodulin-dependent protein kinase IV kinase Isoform [rats, brain, mRNA, 3429 nt]
23474		1.36	8.0E-7		EST_HUMAN	yv46g10.s1 Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:245826 3'
14677	27390	2.59	7.0E-7		LN	Homo sapiens NKG2D gene, exon 10
16082	28732	1.08	7.0E-74		IN	Homo saplens chromosome 21 segment HS21C046
21873	32038	2.83	7.0E-74	BE967432.1	EST_HUMAN	601849284F1 NIH_MGC_73 Homo sapiens cDNA clone IMAGE:3932997 5'
24858	31011	5.61	7.0E-74	BE266305.1	EST HUMAN	601191927F1 NIH_MGC_7 Hamo saplens cDNA clone IMAGE:3535855 5
13858	26518	2.4	6.0E-74		Ę	Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, partial cds
15039	27778	11.78	6.0E-74		L HUMAN	601283521F1 NIH_MGC_44 Hamo sepiens cDNA clone IMAGE:3605453 5'
15039	27777	11.78	6.0E-74	BE388260.1	EST_HUMAN	601283621F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3605453 5'
15634	28279	1.32	6.0E-74		EST_HUMAN	UI-H-BI0-aah-h-03-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA done IMAGE:2709365 3'
15634	28280	1.32	6.0E-74		EST_HUMAN	UFH-BI0-aah-h-03-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:27093653'
16453	29092	1.34	6.0E-74	BE048846.1	EST_HUMAN	hr54e11.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3132332 3'
16453	29093	1.34		BE048846.1	EST_HUMAN	hr54e11.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:3132332.3'
18086	30744	3.02	6.0E-74	11056013	NT	Homo sapiens actin filament associated protein (AFAP), mRNA
13654	26322	7.33	5.0E-74		EST HUMAN	of17c09.y1 Morton Fetal Cochlea Homo sapiens cDNA clone IMAGE:2483704 5'
15413		2.62	. 5.0E-74		EST_HUMAN	PM0-CT0289-271099-001-h07 CT0289 Homo sapiens cDNA
18125	30784	1.86	5.0E-74	11425417	NT	Homo sapiens phosphatidylinositol glycan, class L (PIGL), mRNA
18497	31419	12.98	5.0E-74		LN	H.sapiens mRNA for TPCR16 protein
						Homo sapiens VAMP (vestcle-associated membrane protein)-associated protein A (33kD) (VAPA) mRNA,
18540	31462	7.41	5.0E-74	4507866		and translated products
18608	31536	1.84	5.0E-74	11431471		Homo sapiens interleukin 4 receptor (IL4R), mRNA
18608	31537	1.84	5.0E-74	11431471	NT	Homo saplens interleukin 4 receptor (IL4R), mRNA
19539	32567	5.98	5.0E-74	7662263	IN	Homo sapiens KIAA0716 gene product (KIAA0716), mRNA
19539	32567	9.0	5.0E-74	7662263	NT	Homo sapiens KIAA0716 gene product (KIAA0716), mRNA
20631	33758	2.78	5.0E-74	11345483	LN	Homo sapiens hypothetical protein FLJ13222 (FLJ13222), mRNA
23328	36565	2.56			INT	H.saplens mRNA for HIP-I
23328	36566	2.56			NT	H.sapiens mRNA for HIP-I
23442	36687	2.68	5.0E-74	5729766	Ę	Homo sapiens cell adhesion molecule with homology to L1CAM (close homologue of L1) (CHL1), mRNA
13080	25723	1.79	4.0E-74		FN	Homo sapiens DNA for amyloid precursor protein, complete cds
13602	26272	5.15	4.0E-74		NT	Homo sapiens mRNA for KIAA 1019 protein, partial ods
		ğΩ	ORF SEQ Expres ID NO: Sigr 26150 31644 31644 31644 31646 28732 28732 28279 28279 28279 28279 28279 28279 28279 28279 28279 31636 31636 31636 33758 33758 33567 32567 32567 32567 32567 32567 32567 32567 32567 32567 32567 26272	ORF SEQ Expression (Top) Hit Signal BLAST E Value Value Signal 2.2 8.0E-72 8.0	ORF SEQ Expression Signel (Top) Hit Top Hit Acession Value Top Hit Acession No. 26156 1.06 8.0E-74 4557426 31545 2.2 8.0E-74 S83194.1 31546 2.2 8.0E-74 S83194.1 27390 2.59 7.0E-74 N5239.1 27390 2.59 7.0E-74 AL163246.2 35038 2.83 7.0E-74 BE38280.1 28732 1.08 7.0E-74 BE38280.1 28732 1.08 7.0E-74 BE38280.1 28732 1.08 7.0E-74 BE38280.1 28279 1.32 6.0E-74 BE38280.1 28279 1.32 6.0E-74 BE38280.1 28280 1.34 6.0E-74 BE388280.1 28290 1.34 6.0E-74 BE04834.1 28003 1.34 6.0E-74 AW014039.1 28021 1.34 6.0E-74 AW014039.1 28032 1.34 6.0E-74 AW020988.1 28033 1.34 6.0E-74 AW020988.1 28034 1.28 5.0E-74 AW020988.1 31462 1.84 5.0E-74 AW020	ORF SEQ Expression Signal (Top) Hit Top Hit Acession Signal Top Hit Acession No. Top Hit Acession Signal Top Hit Acession No. Top Hit Acession Signal Top Hit Acession No. Top Hi

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_		_		_		-		_		_	_	_					_		_	_	_	_			_	_	_			_	_		_		
	Top Hit Descriptor	Human platelet glycoprotein IIb mRNA, 3' and	RC6-HT0678-220500-011-C03 HT0678 Homo saplens cDNA	Homo sapiens PDZ-73 protein (PDZ-73/NY-CO-38), mRNA	Homo sapiens PDZ-73 protein (PDZ-73/NY-CO-38), mRNA	Homo saplens PDZ-73 protein (PDZ-73/NY-CO-38), mRNA	Homo sapiens PDZ-73 protein (PDZ-73/NY-CO-38), mRNA	601557524F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3827549 5	Homo saplens mRNA for KIAA1395 protein, partial cds	Homo sapiens chromosome 21 segment HS210004	zp96a06.s1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:628018 3'	Homo sapiens Misshapen/NIK-related knase (MINK), mRNA	QV4-ST0234-181199-037-705 ST0234 Homo sapiens cDNA	Homo sepiens hypothetical protein FLJ11026 (FLJ11026), mRNA	Homo saplens beta 2 gene	Homo saplens zinc finger protein 259 (ZNF259) mRNA	Homo sapiens mRNA for KIAA0833 protein, partial cds	Homo saplens chromosome 21 segment HS21 C046	Homo saplens DNA for Human P2XM, complete cds	Homo saplens mannosidase, alpha, class 2A, member 1 (MAN2A1), mRNA	2r60c01.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:687776 5	zr60c01.r1 Soares_NhHMPu_S1 Homo sepiens cDNA clone IMAGE:667776 5'	Homo sapiens glutamate receptor, Ionotropic, kalnate 1 (GRIK1) mRNA	Homo sapiens glutamate receptor, lonotropic, kalnate 1 (GRIK1) mRNA	Homo saplens chromosome 21 segment HS21C068	RC2-BT0642-270300-019-f06 BT0642 Homo sapiens cDNA	Human neurofibromin (NF1) gene, complete cds	Homo saplens KIAA0852 protein (KIAA0852), mRNA	601070088F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3456260 5'	601070088F1 NIH_MGC_12 Homo sepiens cDNA clone IMAGE:3456260 5'	Homo saplens tracheal epithelium enriched protein (PLUNC) gene, complete cds	MR0-HT0559-230500-021-a03 HT0559 Homo sapiens cDNA	Homo saplens partial AK155 gene for AK155 protein, exons 1-3 and joined CDS	Homo sapiens partial AK155 gene for AK155 protein, exons 1-3 and joined CDS	Homo sapiens hypothetical protein FLJ10783 (FLJ10783), mRNA
,_	Top Hit Database Source	IN	EST_HUMAN	N	LZ LZ	뉟	ᅜ	EST_HUMAN	N		EST_HUMAN	N T	EST_HUMAN	NT	NT	NT	NT	F F	IN	NT	EST_HUMAN	EST_HUMAN	NT	NT	NT	EST_HUMAN	NT	Į,	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	NT	NT	N
	Top Hit Acession No.	.74 J02963.1	74 BE711134.1	11439587 NT	11439587 NT	11439587 NT	11439587 NT	74 BF030788.1	AB037816.1	-74 AL163204.2	.74 AA196181.1	7657334 NT	.74 AW816405.1	8922829 NT	74 X02344.1	4508020 NT	74 AB020640.1	74 AL163246.2	.74 AB002059.1	4758697 NT	74 AA258549.1	74 AA258549.1	4504116 NT	4504116 NT	74 AL163268.2	74 BE083080.1	74 M89914.1	11417977 NT	74 BE549105.1	74 BE549105.1	74 AF214562.1	74 BF351951.1	74 AJ251550.1	74 AJ251550.1	11420549 NT
	Most Similar (Top) Hit BLAST E Value	2.0E-74		2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	2.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74	1.0E-74		1.0E-74		1.0E-74
	Expression Signal	1.11	2.5	1.89	1.89	2.78	2.78	0.92	1.29	90'9	3.95	1.89	5.02	1.05	13.6	1.47	1.81	2.27	4.39	3.55	6.0	6.0	98'0	0.86	4.81	1.15	1.86	1.15	1.13	1.13	4.92	99'0	0.55	0.55	1.38
	ORF SEQ .ID NO:	30286	31424	31521	31522	31521		32748		35208		25509	25764	25907	25912	25995	26197	26406	27690	28545	28782							33291	33773	33774	34543				36241
	Exen SEQ ID NO:	17676		25077	25077	1	25077	19695			24687	12881	13129	13272	13277	13367	13538	13744	14951	16900			16651	16651	16700		19368	20197	20650	L		21425			23026
	Probe SEQ ID NO:	4949	60/9	9089	5806	5876	5876	7003	7841	9282	12234	52	328	487	493	587	765	979	2223	3136	3366	3366	3901	3901	3950	4042	6605	7526	7955	7955	8704	8733	10140	10140	10380

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			Т	T	\top	7		τ		_	_	$\overline{}$	_	_	_	_						_			_				_					
Single Exon Probes Expressed in Brain	Top Hit Descriptor		Homo saplens glutathione S-transferase theta 2 (GSTT2), mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2), mRNA	Homo sapiens DNA for Human P2XM, complete cds	normo septens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	Puro especial Charles	Homo sapilans chamacam 24.	wk38e08.x1 NC_CGAP_P72 Homo sapiens cDNA clone IMAGE;2417654 3' similar to gb;M14123_cds4	HOURUS-RELATED POL POLYPROTEIN (HUMAN);	Intereducts 1 NCI_CGAP_Co3 Homo saplens cDNA clone IMAGE:1028933 31	nkeedus.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:1028833 3'	00 1120J0817 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:2989865 5	2017e0e.r1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:587174 5'	601346505E4 NIT MCC_8 Home sapiens cDNA clone IMAGE:3687458 5	27 134 280 BF I MIN _MCC_8 Homo sapiens cDNA clone IMAGE:3687458 5	#31c12x1 NCI CGAP GC8 Home seniens cDNA clane IMAGE:4298738 3'	HYPOTHETICAL 20.1 KD PROTEIN:	QV1-BT0632-210200-079-e02 BT0632 Homo sapiens cDNIA	W80h08.r1 Scares melanced 2NhHM Homo confine a control	CM0-NN0057-150400-335-811 NN0057 Home series conv. Action (IMAGE: 269055 6)	601303868F1 NIH MGC 21 Homo seniens cDNA close 20170	Homo saplens hypothetical protein FLJ10747 (FLJ10747) mRNA	Homo sapiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1) mRNA	Homo sepiens NIPSNAP, C. elegans, homolog 1 (NIPSNAP1) mRNA	Homo sepiens eukaryotic translation initiation factor 3 submit 8 (140kD) /E12359) - BM4	Homo saplens NIPSNAP, C. elegans, homolog 1 (NIPSNAPA), mBNIA	Homo sepiens NIPSNAP, C. eledens, homology (NIPSNAPA) SPANA	Homo saplens myosin, heavy polypeptide 1. skaletal muscle, edult (AVCLA) Bh.A	Homo saplens HTRA serine prolease (PRSS11) dans complete add	Homo sapiens HTRA serine protease (PRSS11) cena complete colo	Homo sapiens mRNA for KIAA0581 protein partial cds	Homo saplens synaptojanin 1 (SYNJ1), mRNA	Homo sepiens synaptosomal-associated protein, 28kD (SNAP28) mRNA
gle Exon Pro	Top Hit Database Source	1	- N	Z	Ž	¥	Ę	Į.	FOT LIBRAN	Т	Т	EST CINAN	Т	_	EST LIMAN	Τ	Т		Т	Г	HUMAN	EST_HUMAN												
Sin	Top Hit Acession No.			1141/856 N	TA MOOUZOSB. I	-74 AF240786.1	-75 AF178228.1	75 AL163202.2	.75 AI817415 1		T	Ī							1.1			BE409464.1	8922637 NT	11417946 NT	11417946 NT	5579457 NT	11417946 NT	11417946 NT	39505			5 AB011153.1 NT	4507334 NT	4759153 NT
	Most Similar (Top) Hit BLAST E Value	1 05 74	1 05 74	1 05 74	100	1.0E-74	8.0E-75	8.0E-75	6.0E-75	5.0E-75/	5.0F-75	5.0E-75	5.0E-75	5.0E-75		5.0E-75	Г	5.0E-75 A	4.0E-75 B		4.0E-75 A	io [4.0E-75	4.0E-75	4.0E-75	4.0E-/5	4.0E-75	4.0E-75	4.0E-75	3.0E-75 A	3.0E-75 AF	3.0E-75 AE	3.0E-75	3.0E-75
	Expression Signal	200	504	- F.		1.53	3.68	1.86	1.47	0.61	0.61	0.94	0.6	0.78	0.78	1.53		2.39	2.18	1.02	1.5	4.89	0.94	0.56	0.30	0,'0	2.28	2.28	18.12	3.72	2.41	2.76	1.1	5.86
	ORF SEQ ID NO:	37800		27690					27781	33466	33467	34643	34855	34946	34947	35197	-	32883	20208		27202	28266	20802	31120	91036	00000	32430	32437	36517	2040B	26409	2/2/8	27570	6/0/0
	Exon SEQ ID NO:	24458	L	L	1		_	24700	15044	20352	20352	1 1		21782	21782	22027	00200	70,77	1000	13532	14501	200	18244	1824	18052	16/34	70707	7400	E7757	19747	13/4/	7007	15143	2
	Probe SEQ ID NO:	11885	11965	12103		12567	2 <u>8</u> 20	12254	2319	7688	7688	8806	9015	8083	9083	8273	10137	3 5	2 9	\$ 1	1/28	300	5442	5442	8178	9881	age 4	198	5 6	3 8	3 6	1020	2422	6744
																										_	_		_			_		_

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	_	_	_	_	-	_	_	_	_		_		_				_								_	_	_		
Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21C001	Homo saplens mRNA for KIAA0581 protein, partial cds	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds	Human calcium-dependent phospholipid-binding protein (PLA2) mRNA, complete cds	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens KIAA0971 protein (KIAA0971), mRNA	Homo sapiens adaptor-related protein complex 1, sigma 2 subunit (AP1S2), mRNA	Homo sapiens adaptor-related protein complex 1, sigma 2 subunit (AP1S2), mRNA	Homo sapiens HIR (histone cell cycle regulation defective. S. cerevisiae) homolog A (HIRA). mRNA	Homo sapiens HIR (fristone cell cycle regulation defective. S. carevisiae) homology A (HIRA) mRNA	Homo sapiens KIAA0623 dene product (KIAA0623) mRNA	Homo saplens KIAA0623 gene product (KIAA0628) mRNA	Homo sapiens Oncogene TIM (TIM) mRNA	Homo sapiens Oncogene TIM (TIM) mRNA	Homo sapiens snail 1 (drosophila hamdog), zlnc finger protein (SNAI1), mRNA	Homo sapiens Drosophila Kelch like protein (DKELCHL), mRNA	Homo saplens symuclein, alpha (non A4 component of amyloid precursor) (SNCA) mRNA	AV734680 cdA Homo saplens cDNA clone cdABED025	qo91e02.x1 NCI_CGAP_Kid5 Hamo sapiens cDNA clone IMAGE:1915898 3' similar to TR:Q69386 Q69386 POL/ENV GENE	Homo saplens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA	Homo sapiens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA	xg60d02.x1 NCI_CGAP_Ut4 Homo sepiens cDNA clone IMAGE:2632707 3' similar to contains PTR7.tt. PTR7 repetitive element:	H.sapiens ERCC2 gene, exons 1 & 2 (partial)	zt57h03.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:726485 3' similar to gb:M13932 40S	NIBOSOUNAL PROJECT (HOMAN);	601900284F1 NIH MGC 19 Homo sapiens cDNA clone IMAGE:4129678 5	601900294F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4129678 5/	ac77b08.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:868599 3/	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
Top Hit Database Source	N.	LN	N T	L	L _Z	Z	N	NT	Ę	Ę	Ę	ĻΝ	LN FN	TN	L	LN L	N _T	EST_HUMAN	EST_HUMAN	LN	Į.	EST HUMAN	NT	To be	T	T	T	EST_HUMAN	Ę
Top Hit Acession No.	5 AL163201.2	'5 AB011153.1	'5 M72393.1	5 M72393.1	5 D87675.1	7662421	11420956 NT	11420956 NT	11526319 NT	11526319 NT	7662209 NT	7662209 NT	4885632 NT	4885632 NT	11420804 NT	11420222 NT	11436430 NT	AV734680.1	Al311783.1	4506328 NT	4506328 NT	AW 168135.1	X52221.1	A 0300070 4	AA3882/0.1		T	AA664377.1	AF223391.1
Most Similar (Top) Hit BLASTE Value	3.0E-7	3.0E-7	3.0E-7	3.0E-7	3.0E-7	3.0E-7	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	2.0E-75		1.0E-75	1.0E-75	1.0E-75	1.0E-75	100 75	101-101	1.0E-/3		1.0E-75	1.0E-75
Expression Signal	76.0	1.32	0.75	0.75	3.27	0.73	0.83	0.83	1.68	1.68	4.56	4.56	2.52	2.52	121	0.77	2.28	1.41	2.45	1.12	1.12	6.68	3.27	70.1	17.5	0.70	3.79	10.83	2.56
ORF SEQ ID NO:								30494	32623		32783	32784			34717	35414	36314			27539	27540	27762	28366		254.67	33107	00100		36972
SEQ ID NO:								17980	19588		19727	19727			21574	- 1		18384		- 1	14808	15026	15713	21006		21990	- 1		23704
Probe SEQ ID NO:	3021	. 3184	3345	3345	4147	4404	5171	5171	6671	6671	7035	7035	7522	7522	8883	9577	10440	5587	8648	2076	2076	2301	2947	8313	8 8	3 8	2702	16/6/	11033

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Table 4
Single Exon Probes Expressed in

Single Exon Probes Expressed in Brain	Most Similar (Top) Hit Acession (Top) Hit Acession iD NO: Signal BLASTE No. Source Source	37785 1.38 1.0E.75 AAA474434	30598 1.84 1.0E-75 BE8941921 FET HIMAN	25493 1.24 9.0E-76 AI652648.1 EST HUMAN	25494 1.24 9.0E-76 A1652648 1 ECT LINAAN	35654 43.62 9.0E-76 M12937.1 NT	26347 1.06 8.0E-76 4504374 NT	26348 1.06 8.0E-76	28325 1 8.0E-76 7708724 NT	31825 6.36 8.0E-76 1	33145 1.26 8.0E-76 11435215 NT	33231 0.86 8.0E-76 11419212 NT	34027 0.67 8.0E-76 11416981 NT	34835 0.55 8.0E-76 AB046764.1 NT	36137 1.35 8.0E-76 M13792.1	36497 4.81 8.0E-76 10442824 INT	2 8.0E-76 11417862 NT	70786	28607 207 TAT C.DE-70 5016092 NT	28704 7 EE 7 7 70	7.0E-76 4505052NT	28743 0.93 7.0E-76 4757915 NT	29705 4.73 7.0E-76 4507184 NT	29706 4.73 7.0E-76 4507184 NT	31.63 8.0E-76 BE396253.1 FST HIMAAN	38440 3.76 6.0E-76 BE273201.1 EST HUMAN	27385 4.83 6.0E-76 D63874.1 NT	27386 4.83 5.0E-76 D63874.1 NT	27387 4.83 5.0E-76 D63874.1 NT	30512 1.26 4.0E-76 BE783412.1 EST HUMAN	35768 6.42 4.0E-76 D81625.1 EST HUMAN
	ORF SEQ ID NO:	37785	30598	25493	25494	35654	26347	26348	28325	31825	33145	33231	34027	34835	36137	36497		200	28607	28704	10/03	28743	29705	29706		38440	27385	273,86	27387	30512	35768
	SEQ ID NO:	24444		12872			_		_		20067		_1	.	22825	23260	24849	13531	ı	1	1	16090	17077	17071	13962	23209	14671	14671	14671	17886	22571
	Probe SEQ ID NO:	11860	12152	43	43	9801	8	917	2910	6079	388	7466	8196	8995	10277	10564	12491	759	3288	3294		3330	4538	4338	1212	11442	1836	88	1936	2188	8823

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Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9923	22571	35769	6.42	4.0E-7	76 D81625.1	EST_HUMAN	HUM178C01B Human fetal brain (TFujiwara) Homo sapiens cDNA clone GEN-178G01 5'
615		26026		3.0E-7	76 BF516282.1	EST_HUMAN	UI-H-BW1-enz-b-04-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3083862.3
615	13383	26027	3.2	3.0E-7	76 BF516262.1	EST_HUMAN	UI-H-BW 1-anz-b-04-0-UI.s1 NCI_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3083862.3'
1594		27029	8	3.0E-7	4503476 NT	N-	Homo sapiens eukaryotic translation elongation factor 1 beta 2 (EEF1B2) mRNA
1594				2.0€	4503476 NT	Ŋ	Homo saplens eukaryotic translation elongation factor 1 beta 2 (EEF1B2) mRNA
3422					76 BF375689.1	EST_HUMAN	RC5-ST0300-180100-033-A03 ST0300 Homo sapiens cDNA
3422					3.0E-76 BF375689.1	EST HUMAN	RC5-ST0300-180100-033-A03 ST0300 Homo saplens cDNA
4058	16803	29434	1.07	3.0E-76	BE348693.1	EST_HUMAN	ht67112.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMACE:3151823 3' similar to TR:094886 094886 KIAA0792 PROTEIN ;
5158	17891	37796	2.07	3.0E-76	3.0E-76 Z41314.1	EST_HUMAN	HSCZQD042 normalized infant brain cDNA Homo saplens cDNA clone c-zqd04 3'
5646	18441	31354	1.09		3.0E-76 AA160611.1	EST HUMAN	zo/3c07.r1 Stratagene pancreas (#937208) Homo saplens cDNA clone IMAGE:592524 5' similar to gb:L32978 MIXED LINEAGE KINASE 1 (HUMAN):
6275	19048	32025	9.57			Ν	Homo sapiens angiostatin binding protein 1 mRNA, complete cds
8050		33877	0.88	3.0E-7	8 N42671.1	EST_HUMAN	y/20g10.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:271842.5"
9618	22269	35456		3.0E-78	8 AW 299353.1	EST_HUMAN	xs49h01xt NCI_CGAP_Ktd11 Home sapiens cDNA clone IMAGE:2773009 3'
9641		35486	0.99	3.0E-76	6 AA442309.1	EST_HUMAN	zx54d11.11 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:757481 5'
9641	22283			3.0E-76	6 AA442309.1	EST_HUMAN	zv54d11.r1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:757481 5'
11876				3.0E-76	6 AW967984.1	EST_HUMAN	EST380059 MAGE resequences, MAGJ Homo sapiens cDNA
11979			4.86	3.0E-76	'6 AW956455.1	T_HUMAN	EST388525 MAGE resequences, MAGD Homo sapiens cDNA
275		25725	1.59	2.0E-76	6 D84295.1		Human mRNA for possible protein TPRDII, complete cds
333		25768	4.39	2.0E-76	6 D84295.1	· LN	Human mRNA for possible protein TPRDII, complete cds
333		25769	4.39	2:0E-76	6 D84295.1	NT	Human mRNA for possible protein TPRDII, complete cds
448			1.19	2.0E-78	4557662 NT	NT	Homo septens finmunoglobulin (CD79A) binding protein 1 (IGBP1) mRNA
576		25984	1.08	2.0E-78	4503944 NT	NT	Homo sapiens glucagon (GCG) mRNA
1008		26430	1	2.0E-76	4758053 NT	TN	Homo saplens cAMP responsive element binding protein 1 (CREB1) mRNA
1526	14273	26960	1.91	2.0E-76	4504028 NT	FZ	Homo sapiens GM2 ganglioside activator protein (GM2A) mRNA
1526	14273	26961	1.91	2.0E-76	4504028 NT		Homo sapiens GM2 ganglioside activator protein (GM2A) mRNA
1921	14658	27369	0.91	2.0E-78	6 AA253954.1	EST_HUMAN	zs60h11.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:701925 3'
2846	15614	28261	3.34	2.0E-76	6 P23266	SWISSPROT	OLFACTORY RECEPTOR-LIKE PROTEIN F5
3291	16052	28701	2.08	2.0E-76	6 AA445992.1	EST HUMAN	zw64e02.s1 Soares, testis_NHT Homo sapiens oDNA clone IMAGE:780988 3' similar to SW:ITB5_HUMAN P18084 INTEGRIN BETA-5 SUBUNIT PRECURSOR.:
3291	16052	28702	208	2 0E-78		П	ZWG4e02.s1 Scares_tests_NHT Homo sapiens cDNA clone IMAGE:780986 3' similar to SW:ITB5_HUMAN PARA INTEGRIN BETA.s ci ital init poech locop
		l		21.20		1	

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				_	_	_							_	_																	
Single Exon Probes Expressed in Brain	Top Hit Descriptor	ac83b02.y6 Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:869163 5' sImilar to TR:014591	Himan mBNA for sociality and the second seco	OV3-0 Tinny 2 20 and 23 k44 O Towns Li	Homo sanlans EGE like resents and all a life in it.	Gorilla dorilla olfactory recentry (20018) and	Homo sepiens mRNA for KIA 4 1081 model: 2011	Homo sabiens A kinase (PRKA) protect protect 10 (AVABA) Third	Homo saplens TPCR86 profein (HSTPCR88D) - DNA P10, mKNA	Homo saplens similar to ribosomal protein S28 (H contains) (1 Oceases)	Homo sapiens HIRA Interacting protein 4 (Apr. Hills) (LUCoos) 101.	Human mRNA for HMC-1 complete control (Internation), micro A	Human mRNA for HMG-1 complete cds	ROTSEQUEDET NIH MOC 7 Language cus	EST37301 Finhor & week Home content of the state of the	801512435E NILL MCC 74 L	801302333E1 NIH MGC 21 L. Come septemb cone integral 3737 6	yp11402.71 Soarse breast SUBBSH Home septens CDNA clone IMAGE:187165 5' similar to	601866926F1 NIH MGC 17 Home can join 2018 Class at	Homo sapiens professome (prosome, macropain) 28S subunit, non-ATPase, 7 (Mov34 homolog) (PSMD7)	MIKINA 2282007 (Second College Michael III	28526071 Sparse reline N2M-III Dumo Sapiens CUNA cione IMAGE:363578 5	ye89f04.s1 Soares fetal liver spleen 1NFLS Homo sapiens CDNA clone INAGE:123007 3' similar to contains	MENTO repetitive element;	Homo sanians polymerese (PNA N II 7 NA JII 14 14 15 14 15 14 15 14 15 19 19 19 19 19 19 19 19 19 19 19 19 19	Homo saplens polymerase (RNA) II (DNA directed polypeptide E (25kD) (POLKZE) mRNA	Homo septiens interferon (alpha hata and concas) records 2 (151/152). Exist	GP77n12.x1 Soares fetal lung NbH1 19W Homo senions COND Acts 1880 DE 1872 AND DE 1872	7 Homo sapiens diucokinase (GCK) dana awan 2	Homo saplens disintegrin and metalloprotease domain 10 (ADAM10) mRNA	Homo septens tousled-like kinase 1 (TLK1) mRNA, complete cds
gie Exon Pro	Top Hit Database Source	EST HIMAN	L L	FST HIMAN	LV LV	L	Ł	F	님	Ę	Ę	Į	LN	EST HIMAN	EST HUMAN	EST HIMAN	EST HIMAN	FOT LIMAN	EST HUMAN	<u> </u>	T HIMAN	Т		Т				T HUMAN	Г		F
ulo	Top Hit Acession No.	76 AI821149 1	-76 D84295 1	-76 AW879618.1	5031660 NT	AF127845.1		8069		11437211 NT	7549807 NT	76 D63874.1		17		Γ	T		-	0000	2530	8.0E-77 AA019770.1		-	05944	4505944 NT	4504600 NT	7 AI204066.1 E		37250	7 AF162666.1 N
	Most Similar (Top) Hit BLAST E Value	2.0E-76		2.0E-76	2.0E-76	2.0E-76	2.0E-76	2.0E-76	2.0E-76	2.0E-78	2.0E-76	1.0E-76		1.0E-76	1.0E-76	9.0E-77		8 0E-77	8.0E-77	8 AE 77	8.0E-77/	8.0E-77	8.0E-77 R00245 1	7.0E-77	7.0E-77	7.0E-77	. 6.0E-77	6.0E-77	5.0E-77 A	5.0E-77	5.0E-77
	Expression Signal	0.97	1.23	6.21	1.49	1.6	6.47	0.75	1.91	3.53	3.58	2.38	2.38	6.12	0.72	4.53	4.1	0.92	1.09	1 74	191	1.91	4.02	24	2.52	2.52	8.53	3.22	2.11	1.77	0.98
	ORF SEQ ID NO:	28879		L								29638		30849		32508		25633	29849	30854	37351	37352	31008	27370	27868	27869	25703	26969	26631	26763	28139
	Exon SEQ ID NO:	16226		17622	17774			ı		- 1	23521	17006	17006	18164	18927	19486	24941	12894	17221	18168	24048	24048	24925	14659	15132	15132	13064	14281	13964	14087	15400
	Probe SEQ ID NO:	3469	4114	4895	Ш	6226	5531	7334	7560	10182	10839	4265	4265	5362	6150	6825	12644	182	4486	5366	11360	11360	12620	1922	2411	2411	256	1534	1214	1339	2691
																					_			_			_		_		_

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Probe Exon SEQ ID NO: NO: NO: 2767 15472 3512 16288 4655 17389 4884 17611 6886 19603 7730 20393 8266 20966 8266 8266 8266 8266 8266 8266 8266 8	g ₀	Signi		Top Hit Acession No. 4503160 8394516 8394516 5031660 AL043953.1 M13976.1 11428849 11422849	Top Hit Database Source Source NT NT EST_HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	Top Hit Descriptor Homo sapiens cullin 1 (CUL.1) mRNA Homo sapiens cullin 1 (CUL.1) mRNA Homo sapiens biquitin specific protease 18 (USP18), mRNA Homo sapiens EGF-like repeats and discodin Hike domains 3 (EDIL.3), mRNA Homo sapiens EGF-like repeats and discodin Hike domains 3 (EDIL.3), mRNA DKFZp434G1728 r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434G1728 5' Homo sapiens protein kinase C bete-li type (PRKCB1) mRNA, complete cds Homo sapiens hypothetical protein FLJ20343), mRNA Homo sapiens 3-hydroxylsobutyny-Coenzyme A hydrolase (HiBCH), mRNA Homo sapiens 3-hydroxylsobutyny-Coenzyme A hydrolase (HiBCH), mRNA
22078 23034 24384 24384 14701 14701	36250 36249 36250 37716 37717 27417 27418	2.48 1.22 3.12 3.12 3.12 1.09 1.09	6.0E-77 6.0E-77 6.0E-77 5.0E-77 3.0E-77 3.0E-77	5.0E-77	NT NT NT NT NT NT NT EST_HUMAN	Homo saplens sorting nexth 5 (SNX5), mRNA Human mRNA for KIAA0299 gene, partial cds Human mRNA for KIAA0299 gene, partial cds Human UNC-104- and KIF1A-related protein mRNA, partial cds Human UNC-104- and KIF1A-related protein mRNA, partial cds Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA yu64g01.r1 Weizmann Olfactory Epithelium Homo saplens cDNA clone IMAGE:238608 5' similar to SP:S17447 S17447 PROBABLE LIGAND-BINDING PROTEIN RY2G5-:
22836 23468 14079 14160 14816		0.79 3.68 1.4 1.91 1.13	3.0E-77 2.0E-77 2.0E-77 2.0E-77 2.0E-77	2.1	T_HUMAN T_HUMAN T_HUMAN	yu84g01.r1 Weizmann Offactory Epithelium Homo sapiens cDNA clone IMAGE:238608 5' similar to SP:S17447 S17447 PROBABLE LIGAND-BINDING PROTEIN RY2G5 -; PM3-MT0078-080300-005-g03 MT0078 Homo sapiens cDNA AV764617 MDS Homo sapiens cDNA clone MDSBTF10 5' RC3-BN0053-170200-011-h01 BN0053 Homo sapiens cDNA Homo sapiens CYP17 gene, 5' end Homo sapiens CG-79 protein (LOC51634), mRNA
15599 15599 16758 17116	28053 28054 29386 29749 29750	1.96	2.0E-77 2.0E-77 2.0E-77 2.0E-77	2.0E-77 AB037836.1 II 2.0E-77 AB037836.1 II 2.0E-77 AB04316.1 II 2.0E-77 AI613519.1 II	NT EST_HUMAN EST_HUMAN EST_HUMAN	Homo saplens mRNA for KIAA1415 protein, partial cds Homo saplens mRNA for KIAA1415 protein, partial cds Homo saplens mRNA for KIAA1415 protein, partial cds ho43b05.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:3040113 3' similar to SW:GAG2_HUMAN P10264 RETROVIRUS-RELATED GAG POLYPROTEIN; tw22g02.x1 NG_CGAP_Bm52 Homo sapiens cDNA clone IMAGE:2260466 3' similar to TR:O65245 w22g02.x1 NG_CGAP_Bm52 Homo sapiens cDNA clone IMAGE:2260466 3' similar to TR:O65245 C065245 F21E10.7 PROTEIN;

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		otransferase 2) (GOT2),		THE HOMAN		to TR:013311	D-F29D11 1								mRNA		YNN H	mRNA.	ANA									Ī
Gringle Exploses Exploseed in Brain	Top Hit Descriptor	Homo sapiens glutamic-oxaloacetic transaminase 2, milochondrial (aspartate aminotransferase 2) (GOT2), nuclear gene encoding milochondrial peries. mBNA	ns68g12.s1 NCI_CGAP_Pr2 Homo saplens cDNA clone IMAGE-1188838 cimilize to the control of the con	P47914 60S RIBOSOMAL PROTEIN L29. [1] ;contains element MSR1 repetitive element	601119852F1 NIH MGC_17 Homo saplens cDNA clone IMAGE:3029436 5'	SOLVINOSZET NIT MCC. 88 Hamo saplens cDNA clone IMAGE:3878505 6' MAGE:2877720 5' similar to TR:013311	13 (2009) 1 CAL-BINDING FROI EIN 1 (XBP151, [1]); (2009) 1 CAP_BINZ5 Homo saplens cDNA clone IMAGE:2017380 3 similar to WP-E280111 1	CEUS/69 LOW DENSITY LIPID RECEPTOR-RICATED PROTEIN;	Human protein kinase C substrate 80K-H (PRKCSH) gene, exon 7	Pruman protein kinase C substrate 80K-H (PRKCSH) gene, exon 7	601685183F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4124541 51	Uses 1637 I MIN MAC 19 Homo Sepiens oDNA clone IMAGE:4124641 6/	Homo suprens mKNA for KIAA1276 protein, partial ods	ricino septens rimana for NAA1276 protein, partial cos	Homo sapiens amylold beta (A4) precursor protein (protease nexin-II, Alzhelmer disease) (APP), mRNA	Homo sapiens amyloid beta (A4) precursor protein (protease nextn.!! Alzheimer diseas) (ABD)	OMO caniane amulaid between 18.11.	ment of the same of the production of the process of the same of t	Homo sapiens amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP) mRNA	wwsge05.x1 Scares_thymus_NHFTh Homo saplens oDNA clone IMAGE:2538160 3	right of sapiens miking for KIAA1101 protein, complete cds	Tromo sapiens 2,4-dienoy CoA reductase 1, mitochondrial (DECR1), mRNA	rigino sapiens CGI-60 protein (LOC51626), mRNA	From Saplens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Fromo sapiens breast cancer 1, early onset (BRCA1), transcript variant BRCA1-exon4, mRNA	Home sapiens KIAA0005 gene product (KIAA0005), mRNA	Hours saprens NA40005 gene product (KIAA0005), mRNA Homo sapiens dynactin 1 (DCTN1) rems common and an analysis	capteris unitactiff (CCTINT) gene, exons 27 and 28
	Top Hit Database Source	FN		EST HUMAN	EST HIMAN	EST LINAM	I	NAMOL	Z	101011	T	Т								FO HOMAN								
5	Top Hit Acession No.	4504088 NT		77 BE 208040 4	77 BE787143.1			1			T	T	T		4502166 NT	4502166 NT	4502168 NT	AEDOROGA	7 AMOE9440 4	T	3300	7708299 NT	A.1229041 1	2222	7841840 NT	7661849INT	AF086944.1 NT]
Moet Similar		2.0E-77	Ļ		2.0E-77	2.0E-77	2.0E-77		2.0E-77	2.0E-77.E	2.0E-77	1.0E-77.	1.0E-77	, 1	1,-30-1	1.0E-77	1.0E-77	1 05-77	1 0F-77 A	1.0E-77		1.0E-77	1	1.0E-77	1.0E-77	1.0E-77		
	Expression Signal	0.96	4	1.08	1.73	15.45	0.82	5.05	5.05	0.47	0.47	1.03	1.03	7 40		7.19	17.31	17.34	a o	1.32	1.72	3.37	16.41	2.29	-	-	2.45	
	ORF SEQ ID NO:		30082		31826	32829	34259	35280	35281	35738	35739	25489	25490	25714		25715	26296	26297	27360	27902	28451	29684	29851	29967	30337	30338	31561	
	SEQ ID NO:	17292	17449	L	18859	19765	21120	22106	22108	22545	22545	12870	12870	13074		13074	15554	15554	14649	15164	15806	17059	17223	17338	17732	17732	18627	
4	SEQ ID	4557	4717	5865	9080	7074	8427	9428	9428	9895	9895	42	42	798		997	855	855	1912	2445	3040	4320	4488	4803	5010	6010	6839	

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Table 4
Single Exon Probes Expressed in Brain

	Π	Π	Γ	Γ	Γ		Γ	Γ		Γ		Γ	Γ	Γ	Γ	Γ			T	1	Π	Γ	Γ	Γ		Γ	Π	\neg			П	١,	П
Top Hit Descriptor	Homo saplens dynactin 1 (DCTN1) gene, exons 27 and 28	Human von Willebrand factor gene, exon 20	Homo sapiens diaphanous (Drosophila, homolog) 1 (DIAPH1), mRNA	Homo saplens elastin (supravalvular aortic stenosis, Williams-Beuren syndrome) (ELN), mRNA	Homo saplens cullin 1 (CUL1), mRNA	Human mRNA for kidney epidermal growth factor (EGF) precursor	H.saplens DNA for Cone cGMP-PDE gene	H.saplens DNA for Cone cGMP-PDE gene	Homo saplens hu-GlcAT-P mRNA for glucuronyltransferase, complete cds	Homo saplens hu-GlcAT-P mRNA for glucuronyltransferase, complete cds	RC3-CT0254-280999-011-b05 CT0254 Homo sapiens cDNA	RC2-ET0023-080500-012-e05 ET0023 Homo saplens cDNA	RC2-ET0023-080500-012-e05 ET0023 Homo saplens cDNA	AU118789 HEMBA1 Homo sapiens cDNA clone HEMBA1004354 5'	AU118789 HEMBA1 Homo sapiens cDNA clone HEMBA1004354 5'	Homo saplens GDNF family receptor alpha 1 (GFRA1), mRNA	Homo sapiens hypothetical protein FLJ11316 (FLJ11316), mRNA	ba54h03.y3 NIH_MGC_10 Homo saplens cDNA clone IMAGE:2900405 5' similar to WP:Y48B8A.8 (CE22121 :	Human collagenase type IV (CLG4) gene, exon 6	Homo sapiens Best's macular dystrophy related protein mRNA, partial cds	Homo saplens transforming growth factor, beta-induced, 68kD (TGFBI), mRNA	EST365190 MAGE resequences, MAGB Homo sapiens cDNA	Human lysosomal alpha-mannosidase (manB) gene, exon 7	601648061F1 NIH_MGC_62 Hamo saplens cDNA clone IMAGE:3931887 5	DKFZp434N0323_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N0323 5'	Novel human gene mapping to chomosome 22	WIB7b12.X1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2495615 3' similar to SW:WAP_PIG	O40055 WHEY ACIDIC PROTEIN PRECURSOR;	nomo sapiens pre-mknA splicing factor (SFRS3) mknA, complete cds	Hamo sepiens syncytin (LOC30816), mRNA	Homo saplens phosphatidylinositol 4-kinase, catalydc, alpha polypeptide (PIK4CA) mRNA	Homo sapiens phosphatidylinositol 4-kinase, catalytic, alpha polypeptide (PIK4CA) mRNA	Homo saplens SFRS protein kinase 2 (SRPK2), mRNA
Top Hit Database Source	N	TN	NT	님	FN	N	NT	Ŋ	뒫	시	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	FA.	TN	EST HUMAN	TN	LN	N FA	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	NT	1	EST HUMAN	Z	Ę	Z,	卜	¥
Top Hit Acession No.	AF086944.1	-77 M25844.1	4885182 NT	5881412 NT	11420159 NT	.77 X04571.1	77 X94354.1	77 X94354.1		77 AB029396.1	78 AW753302.1	78 AW947061.1	78 AW947061.1	78 AU118789.1	78 AU118789.1	11432710 NT	11422486 NT	78 AW673424.1		78 AF038536.1	11416585 NT	0.1	78 U60889.1	78 BE960836.1		78 AL355841.1		Ī	/8 AFTU/405.1	7656876 NT	4505806 NT	4505806 NT	11420732 NT
Most Similar (Top) Hit BLAST E Value	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77		1.0E-77	9.0E-78	8.0E-78	8.0E-78	6.0E-78	6.0E-78	6.0E-78	5.0E-78	5.0E-78		6.0E-78	5.0E-78			5.0E-78	4.0E-78	4.0E-78	Lo		4.0E-78		4.0E-78	4.0E-78	4.0E-78
Expression Signal	2.45	1.4	0.82	21.7	1.05	0.69	0.62	0.62	3.1	3.1	2.55	3.11	3.11	1.87	1.87	2.54	0.72	5.53	3.81	2.33	11.12	2.23	6.78	3.31	1.07	1.78		-0.E	Z.08	1.73	2.61	2.61	1.41
ORF SEQ ID NO:	31562	31697			33338					36289		32117			25549		25663	28019	١.				34821	34822	26531	26940	OFATA	6/0/2	81117	29652	30088	30089	31391
Exen SEQ ID NO:	18627				20234					23067					12910	19232	13024	16281	l		18287		21671	21672	13872	14254	7,000	14390	1904	17027	17454	17454	18474
Probe SEQ ID NO:	5839	5956	6356	6953	7564	7663	9165	9165	10421	10421	10449	6354	6354	22	84	6465	212	2567	3380	5327	5488	7054	8981	8982	1115	1508	707	1 0	013	4288	4722	4722	5681

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D SEQ ID ID NO. Signal R.AST E No. Source	obes Expressed in Brain			Top Hit Descriptor		601482143F1 NIH_MGC_68 Homo sapiens cONA close MACE: 388,455,7;	601482143F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE-3884664 E	Homo saplens netrin 1 (NTN1), mRNA	Homo sapiens netrin 1 (NTN1), mRNA	601112055F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3382888 gr	Homo sapiens mRNA for KIAA0620 protein, partial cds	Homo sapiens mRNA for KIAA0620 protein, partial cds	Homo sapiens guanine nucleotide exchange factor for Rap1 (KIAA02771 mBNA	Homo sapiens Bci-2-associated transcription factor short form mBNA	Homo sapiens tetratricopeptide repeat domain 3 (TTC3), mRNA	AV698115 GKC Homo saplens cDNA clone GKCAHF11 F	Homo saplens Bcl-2-associated transcription factor short form mRNA completed.	Homo sapiens Bci-2-associated transcription factor short form mRNA complete 3	yr48f03.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA close MASCE Socretion	601159415F2 NIH_MGC_63 Homo sapiens cDNA clone IMAGE: 2411177 #	Homo saplens BCL2-like 2 (BCL2L2) mRNA	th18h07.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE-211se88# 2/	Homo sapiens Dickkopf gene 4 (DKK-4), mRNA	Homo sapiens Dickkopf gene 4 (DKK-4), mRNA	Homo saplens phosphodiesterase 6A, cGMP-specific and alpha (BDEsA)	Homo saplens phosphodiesterase 6A, cGMP-specific rod slohe (PDEBA) TENIA	Homo sapiens mRNA for Fas-associated factor, FAF1 (Faf1 gens)	forno sapiens mRNA for KIAA0937 protein, partial cds	fomo saplens chloride channel CLC4 (CIC4) mRNA, complete cds	10mo saplens mRNA for Fas-associated factor, FAF1 (Faf1 gene)	ST182926 Jurket T-cells VI Homo sapiens cDNA 5' and similar to Similar to C. elegans hypothetical and the same and the sam	Source and a source of the sou	ionio sapiens X transporter protein 3 (XT3), mRNA	idnio sapiens mRNA for KIAA0830 protein, partial cds	omo saplens membrane-associated calcium-indanam-dent attaches in	отно saplens Rho GTPase activating protein 8 / ARHCAPRI 1000 saplens Rho GTPase activating brotein 8 / ARHCAPRI 1000 saplens	omo saplens Rho GTPase activating protein 6 (ARHCAPA) transcript variant 4, mRNA	Homo sepiens retinoblastome-like 1 (p107) (RBI 1) - BANA
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SEQ ID DRI D			Expression	Bugio	0 0	0 83	3.56	3.56	0.67	3.35	3.35	0.76	1.61	133	124	1.52	1.52	0.99	1.6	228	2.09	1.12	1.12	5.93	583	1.07	1.09	0,83	1.09		1.22	0.8	1.14	- 6	0.89	1,1	108	1,55
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				ne sector																SLC7AB),	SLC7A8),	partial cds					UEM_HUMAN	
Top Hit Descriptor	Homo saplens hypothetical protein FLJ11008 (FLJ11008), mRNA	Homo sapiens hypothetical protein FLJ20275 (FLJ20275), mRNA	Homo sapiens hypothetical protein FLJ20275 (FLJ20275), mRNA	Homo sepiens similar to ATPase, H+ transporting, lysosomal (vacuolar proton pump) membrane sector	descended protein Mode (n. Sapiens) (LOCOSEO), mRNA	HA/D10S170 Emittetive cytestated brother frumes through a DNA 2011 ftt	Human contactin 1 precursor (CNTN1) mRNA complete cde	RC4-BT0310-110300-015-f10 BT0310 Homo seniens cDNA	RC4-BT0310-110300-015-f10 BT0310 Homo sapiens cDNA	Homo saplens KIAA0879 protein (KIAA0879), mRNA	Homo sepiens mRNA for KIAA0833 protein, partial cds	Homo sapiens cadherin EGF LAG seven-pass G-type receptor 1 (CELSR1), mRNA	MR0-NN0087-260600-017-510 NN0087 Homo sapiens cDNA	801311517F1 NIH MGC 44 Homo sapiens cDNA clone IMAGE:3632909 5	QV2-HT0540-120900-358-e05 HT0540 Homo sapiens cDNA	ar 79a04.x1 Barstead colon HPLRB7 Homo sapiens cDNA clone IMAGE:21514383'	al23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	ai23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	601581652F1 NIH_MGC_7 Homo sepiens cDNA clane IMAGE:3836061 5'	Homo sapiens solute carrier family 7 (catlonic amino acid transporter, y+ system), member 8 (SLC7A8), mRNA	Homo saplens solute carrier family 7 (cationic amino acid transporter, y+ system), member 8 (SLC7A8), mRNA	Homo sapiens Y chromosome spermatogenesis candidate protein (RBM) pseudogene mRNA, partal cds	Homo sapiens KIAA0724 gene product (KIAA0724), mRNA	Homo sapiens KIAA0724 gene product (KIAA0724), mRNA	Homo sapiens triple functional domain (PTPRF interacting) (TRIO), mRNA	Homo sapiens triple functional domain (PTPRF Interacting) (TRIO), mRNA	#F68d02.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2103459 3' similar to SW:NUEM_HUMAN Q16795 NADH-UBIQUINONE OXIDOREDUCTASE 39 KD SUBLINIT PRECURSOR:	Homo sapiens NRD convertase mRNA, complete cds
Top Hit Database Source	IN	NT	N	Ŀ	FIA	L L	N.	EST HUMAN	EST HUMAN	Z	N	FZ	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	LZ LZ	L L	N	LN	F	Z-L	FZ	EST HUMAN	۲
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Most Similar (Top) Hit BLAST E Value	2.0E-79	2.0E-79	2.0E-79	07 30 6	20.0	2.0E-79	2.0E-79	2.0E-79	2.0E-79	2.0E	2.0E-	2.0E-79	1.0E-79	1.0E-79	1.0E-79	1.0E-79	9.0E-80	9.0E-80	9.0E-80	9.0E-80	9.0E-80	8.0E-80	8.0E-80	8.0E-80	8.0E-80	8.0E-80	6.0E-80	8.0E-80
Expression Signal	2.26	0.58	0.58	1 22	24.	1 44	1234	4.05	4.05	2.18	5.19	2.89	3.76	0.78	2.05	1.8	2.35	2.35	1.14	8.66	8.66	1.31	3.07	3.07	1.13	1.13	1.12	2.22
ORF SEQ ID NO:			34504	75476	L							31075		33969	37543		28551	28552	35757	37199	37200		33268	33269	35142	35143	26318	27071
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Probe SEQ ID NO:	8415	8664	8864	Ronz	600	6882	10698	10956	10958	11936	12018	12238	6492	8143	11623	12047	3143	3143	9912	11245	11245	3588	7504	7504	8302	9302	880	1638

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B Expn NO: CRF SEQ Signal Expression (Top) Hit Value Top Hit Acess No. Top Hit Acess Value Top Hit Acess No. 1 6993 29620 1.08 6.0E-80 AB032281.1 2 18505 31427 1.79 6.0E-80 AB032281.1 1 18505 31427 1.79 6.0E-80 AJ404468.1 1 18505 31427 1.79 6.0E-80 AJ404468.1 1 18506 31729 3.37 6.0E-80 AJ404468.1 1 18506 31729 3.22 6.0E-80 AJ404468.1 2 18506 31729 3.22 6.0E-80 AJ404468.1 3 1850 0.96 6.0E-80 AJ404468.1 11528.3 3 21416 34559 3.22 6.0E-80 AJ404468.1 3 21416 34559 3.22 6.0E-80 AJ404468.1 2 21638 3.21 6.0E-80 AJ404468.1 3 21416 34559 3.22 6.0E-80 AJ41465.1 3 3773 1.59 6.0E-80 AJ41465.1 3 3772 1.59 6.0E-80 AJ422197.1 2 4392 37726 2.26 6.0E-80 AJ422197.1	gle Exon Pro	Top Hit Database Source		Į.	Į.	I NI	ž į	ž	ž	LN.	Į.	ž!	Į			7			<u> </u>		<u></u>				NAMOL												
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SEC DE DE				1.08	1.08	1.79	337	4 69	1 17	0.98	3.22	3.22	1.51	0.83	1 48	283	28 56	200	60.	1.5	2.26	2.26		1.88		2.42	0/3	\$A.		L		1 07	65.0	2.38	1 89	1.65	
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		Probe SEQ ID NO:	4262	7505	4252	5712	5984	6135	6179	6230	8723	8723	8917	9259	9761	10861	11187	11702	11758	11803	14000	2007		11906	12028	12219	12707		12804	574	818	818	1166	1439	2361		

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Γ		Г	Γ	Π	Γ	Г	Т	Γ	Г	Г	Τ	Т	Т	Τ	ī	Г	Г	Т	Т	Γ^{-}	Γ	Г	œ	T	Т	Г	_		П	7	\neg
	Top Hit Descriptor	Homo saplens H3 histone family, member J (H3FJ) mRNA	Homo sapiens HMT-1 mRNA for beta-1,4 mannosyltransferase, complete cds	Homo sapiens HMT-1 mRNA for beta-1,4 mannosyltransferase, complete cds	Homo sapiens chromosome 21 segment HS21C088	Mus musculus keratin complex 2, gene 6g (Krt2-6g), mRNA	HSPD13155 HM3 Homo saplens cDNA clone s4000045F03	Homo sapiens chromosome 21 segment HS21C010	PM0-GN0018-040900-002-E03 GN0018 Homo sapiens cDNA	QV4-BN0263-040600-241-910 BN0263 Homo saplens cDNA	0023e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:1567054 3' similar to TR-035790 035790 PIG.	vd65a08.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:38060 5'	RET487 subtracted retina cDNA library Homo sapiens cDNA clone RET487	DKFZp434D1323 r1 434 (synonym: htes3) Homo septens cDNA clone DKFZp434D1323 5	nn80d01.s1 NCI_CGAP_Co9 Homo saplens cDNA clone IMAGE:10901773'	Homo saplens Golgi transport complex protein (90 kDa) (GTC90), mRNA	ye86f12.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:22851 6' similar to SP:K1CR, XENLA P08802 KERATIN, TYPE I CYTOSKELETAL ENDO B;	EST378343 MAGE resequences, MAGH Homo sapiens cDNA	Homo sapiens GGT gene, exon 6	zt70f12.r1 Spares_testis_NHT Homo saplens cDNA clone IMAGE:727727 5' similar to TR:G191315 G191313 ANDROGEN-DEPENDENT EXPRESSED PROTEIN.;	Hamo saplens chromosome 21 segment HS21C103	Homo sapiens chromosome 21 unknown mRNA	nn01f12.x5 NCI_CGAP_Co9 Homo saplens cDNA clone IMAGE:1076495 3' similar to contains OFR.tl OFR payelflue element	Homo septiens damma-amilnobutyric acid (GABA) A receptor, gamma 2 (GABRG2) mRNA	601274305F1 NIH_MGC_20 Hamo saplens cDNA clone IMAGE:3615433 5'	Human pro-alpha1 type II collagen (COL2A1) gene exons 1-54, complete cds	Homo sapiens malate dehydrogenase 2, NAD (mitochondrial) (MDH2), nuclear gene encoding mitochondrial	protein, mRNA	Homo sapiens mRNA for lipophilin B	wq25c05.x1 NCI_CGAP_Kld11 Homo sapiens cDNA clone IMAGE:2472296 3'	wq25c05x1 NCI_CGAP_Kid11 Homo saplens cDNA clone IMAGE:2472296 3'
	Top Hit Database Source	TN	N	Ι	ΙZ	N	EST_HUMAN	N	EST_HUMAN	EST_HUMAN	NAMI H TAR	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	TN	EST HUMAN	EST HUMAN	NT	EST_HUMAN	NT	LN	ECT HIMAN	4567610 NT	EST HUMAN	NT.		NT	LZ	EST HUMAN	EST_HUMAN
	Top Hit Acession No.	4504292 NT	30 AB019038.1	30 AB019038.1	30 AL163268.2	9910293 NT	F25915.1	AL163210.2	3.0E-80 BF085009.1	BE817465.1	20 A1091675 1	2.0E-80 R35321.1	2.0E-80 A 444821.1	2.0E-80 At 043116.2	4A582952.1	2.0E-80 11421930 NT	175215.1	4W964270.1	2.0E-80 AJ007379.1	4A393362.1	1.0E-80 AL163303.2	4F231920.1	N 1730858 4	1.0E-80 4567610		1.0E-80 L10347.1		5174540 NT	1.0E-80 AJ224172.1	1.0E-80 AI948731.1	AI948731.1
	Most Similar (Top) Hit BLAST E Value	5.0E-80	5.0E-80	5.0E-80	5.0E-80	5.0E-80	4.0E-80 F25915.1	3.0E-80	3.0E-80	3.0E-80	3.05.80	2.0E-80	2.0E-80	2.0E-80	2.0E-80	2.0E-80	2.0E-80 T75215.1	2.0E-80	2.0E-80	2.0E-80	1.0E-80	1.0E-80 /	1 05 80	1 05-80	1.05-80	1.0E-80		1.0E-80	1.05-80	1.0E-80	1.05-80/
	Expression Signal	1.67	1.37	1.37	1.28	1.04	8.77	8.96	1.7	3.77	2,68	5.08	1.19	5.82	0.93	1.71	1.46	1.41	1	4.49	2.25	1.37	24.4	800	6.43	6.58		1.36	0.95	2.53	2.53
	ORF SEQ ID NO:	28242	29393	29394	30244	34086	35056		30030		31443	27238	27307	27509	32667	32496	32908	34905	36516	36705		26215		30397		31608		32174		33237	İ
	Exen SEQ ID NO:	15502	16764	16764	L	20949	21888	13023	17395	17580	18522	14530	14591	14782		ŀ	ŀ	Ι.		1	13132	13554	14882	1		ı	ı	19175	19794	20145	20145
	Probe SEQ ID NO:	2797	4018	4018	4900	8255	9157	211	4661	4850	6720	1790	1853	2049	6708	6813	7151	9057	8996	10780	331	782	1047	5060	5244	5881		6406	7106	7472	7472

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Most Similar			_	_	_			_		_	_		_	_																						
Exon ORF SEQ Expression Top Hit Acession	bes Expressed in Brain	Top Hit Descriptor		Homo sapiens protein tyrosine phosphatase, receptor type. A (PTPRA) mBNA	Homo sepiens protein tyrosine phosphatase, receptor type. A (PTPRA)RNA	Homo saplens protein tyrosine phosphatase, receptor tyre 4 (BTBA) - BAA	Homo sapiens probable mannose binding Characher 1970; Calain Paris	Homo saplens probable manners himle C the Letter Complete cds	Homo saplens mRNA for KIA And 45 matter.	Homo saplens similar to rat myomoralis (1 Ocassos) — Davis	Home canions cimilar to the manufacture of the manu	Homo saniens marindome (Airman Consequente Marindome (Airman Airman Consequente Marindome (Airman Airman)	Home capters in remitigated tursupted in balanced translocation) 1 (MN1), mRNA	Cholone of Secondary Tor KIAA0833 protein, partial cds	rymogous is owned NPL i GBC S1 Homo saplens cDNA clone IMAGE:1854296 3'	anagous a varies NFL : GBC S1 Homo sapiens cDNA clone IMAGE: 1854296 3	201310331711VIH_MGC_44 Homb saplens cDNA clone IMAGE;3632070 5	SO 11100-XX SOBRES TERM LING NOHL 19W Home sapiens cDNA clone IMAGE: 299918 31	30414102051 NIH_MGC_18 Homo saplens oDNA clone IMAGE:3352840 5	COLUMN MINITARY TO HOMO sepiens cDNA clone IMAGE:3352840 67	Torring septems A I P-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	John Septems A I P-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	Septemble of the complete of t	20109128 Fetal lung II from Sapiens cDNA 5' end	20 CLESZEI FINIT MOC 44 HOMO Saplens CDNA clone IMAGE:3659284 6	102153A66E1 MILL MOC 93 U.	101125505E1 NIH MOO 6 U	Idmo saplens mRNA for KIA AAASA	form contains mENA for KIAAATT	uman transforming and the control of	impart removement grown recorded (with beta) mRNA, complete eds	renneau tenisionning growth teator-beta (tgf-beta) mRNA, complete cds	omo saptens hypometical protein (FLJ11045), mRNA	isolatis amustilio repeat gene deletes in velocardiofacial syndrome (ARVCF), mRNA isolatis versional and construction of the c	DESCRIPTION OF THE PROPERTY OF	Infection NOT COAP CO14 Homo saplens cDNA clone IMAGE:3035907 3' similar to SW:COPG_BOVIN P53620 COATOMER GAMMA SUBUNIT:
e Exon CRF SEQ Expression (Top) Hit Top Hit Acess NO: ID NO: Signal (Top) Hit Top Hit Acess NO: ID NO: Signal (Top) Hit Top Hit Acess 20824 33960 2.84 1.0E-80 11421 11 21283 34436 1.72 1.0E-80 11421 12 21285 34436 1.21 1.0E-80 11421 11 21283 34436 1.21 1.0E-80 AF245219.1 21855 35019 1.21 1.0E-80 AF245219.1 21856 35019 1.21 1.0E-80 AF245219.1 22872 36480 5.25 1.0E-80 AF245219.1 23278 36516 1.21 1.0E-80 AF245219.1 23278 36516 1.46 8.0E-81 BE36823.1 1.14172 24853 31034 3.08 1.0E-80 AF245715.1 1 17082 29726 5.26 1.0E-80 AF3618 1 17082 29726	ngle Exon Pro		1	N	INT	N-I	NT	TN	PA	LN LN	μ	LN	L'N	EST HIMAN	ENT CINANI	FOT LIMAN	EST HIMAN	EST HIMAN	Т	NOW!	NT		T HIMAN	1	Т	Т	Т								1	EST HUMAN P
Exon ORF SEQ Expression Top)	7						AF245219.1	AF245219.1	D63479.2				AB020640.1	Al251752.1	A1251752 1	BE394525 1	AI822115.1	BE256829 1	3E256829.1	450184R	4501848	AF038660 1	T	1		Γ			Γ	Γ		3508R34	11526341 N		T	
Exon ORF SEQ Express				0-10-10-10-10-10-10-10-10-10-10-10-10-10	7.0E-80	1.0E-80	1.05-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80	1.0E-80	8.0E-81	8.0E-81	8.0E-81	7.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	6.0E-81	5.0E-81	5.0E-81	5.0E-81	5.0E-81 N	5.0E-81 N	5.0E-81	5.0E-81	30 00	4.0E-61	4.0E-81 A
Decorption Colored C		Expression Signal	284	1.01	172	7/7	1.21	1.21	0.95	5.25	5.25	1.57	3.08	1.48	1.46	8.46	3.58	5.28	5.28	2.1	2.1	76.0	1.36	1.81	2.29	2.29	2.8	1.83	1.83	0.77	0.77	2.23	1.3		30.7	1.31
8 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0											ł		31034	36515	36516	37048	32909	29726	29727	30830	30831	33253	34989	37723	31041	31042	27682	34143	34144	35383	35384	37491	37764	28109	-	27270
Probe SEQ ID NO: D 8601 8601 8601 8601 8601 8601 8601 12289 112289 110548 110583 11102 7489 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 12430 1658 1658 1658 1658 1658 1658 1658 1658				L	L	L				⊥	L	$_{ m I}$	_[_	23278	23772	19839	17092	17092	18009	18009	20161	21824	24390	24803	24803	14942	21005	21005	22201	22201	24178	24423	13481	-	14555
		Probe SEQ ID NO:	8130	8601	8601	9185	0185	3000	10220	10040	9	12289	12498	10583	10583	11102	7162	4364	4354	5201	2201	7489	9138	11800	12430	12430	22.14	2 2	123	9548	8248	11577	11839	989		1815

Page 379 of 536 Table 4 Single Exon Probes Expressed in Brain

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Top Hit Descriptor	Homo sapiens mRNA for KIAA1345 protein, partial cds	ws90h03.x1 NCI_CGAP_Co3 Hamo saplens cDNA clone IMAGE:2505269 3' similar to TR:043815 043815 STRIATIN	Homo sepiens rab3 interacting protein variant 2 mRNA, partial cds	Homo sepiens rab3 interacting protein variant 2 mRNA, partial cds	Homo saplens hypothetical protein FLJ20220 (FLJ20220), mRNA	Homo sapiens calcium channel, voltage-dependent, L type, alpha 2/delta subunit (CACNA2) mRNA	Homo saplens ets variant gene 1 (ETV1), mRNA	Human mRNA for amyloid A4(751) protein	Human cone photoreceptor cGMP-phosphodiesterase alpha' subunit gene, exons 2 and 3	Human cone photoreceptor cGMP-phosphodiesterase alpha' subunit gene, exons 2 and 3	Homo sapiens mRNA for Death-associated protein kinase 2, complete cds	Homo sapiens ligase I, DNA, ATP-dependent (LIG1), mRNA	Homo sapiens acyl-Coenzyme A dehydrogenase family, member 8 (ACAD8), mRNA	Homo sapiens acyl-Coenzyme A dehydrogenase family, member 8 (ACAD8), mRNA	Homo sapiens vesicle trafficking protein sec22b (SEC22B) mRNA	Homo saplens vesicle trafficking protein sec22b (SEC22B) mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens beta-ureidoproplonase (LOC51733), mRNA	Homo sapiens beta-ureidoproplonase (LOC51733), mRNA	Homo saplens transcobalamin II; macrocytic anemia (TON2), mRNA	Homo sapiens NF2 gene	Homo sapiens NF2 gene	Homo sapiens cullin 4A (CUL4A) mRNA, complete cds	Homo sapiens pleiotrophin (heparin binding growth factor 8, neurite growth-promoting factor 1) (PTN) mRNA	Homo sapiens pleiotrophin (heparin binding growth factor 8, neurite growth-promoting factor 1) (PTN) mRNA	601474072F1 NIH MGC 68 Homo sapiens cDNA clone IMAGE:3877121 5	601474072F1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3877121 5	hg85c01.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE.2952384 3'	Homo sapiens hypothetical protein (LOC55586), mRNA	
Top Hit Database Source	LN	EST_HUMAN	NT	HN	NT	Į.	Ę	LN	LN	TN	NT	NT	F	LN	N.	Ę	Ę	Ę	۲	ΝΤ	TN	NT	NT	NT	F.	L	EST HUMAN	EST HUMAN	EST HUMAN	TN	
Top Hit Acession No.	AB037766.1	4.0E-81 AW004608.1	4.0E-81 AF263306.1	4.0E-81 AF283306.1	8923209 NT	4757893 NT	11420544 NT	4.0E-81 X06989.1	4.0E-81 U20197.1		4.0E-81 AB018001.1	11425281 NT	11439065 NT	11439065 NT	4759085 NT	4759085 NT	11417862 NT	11417862 NT	11417871 NT	11417871 NT	11417974 NT		3.0E-81 Y18000.1		4506280 NT	4506280 NT	2.0E-81 BE784636.1		611542.1	8923839 NT	
Most Similar (Top) Hit BLAST E Value	4.0E-81 AB	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	4.0E-81	3.0E-81 Y1	3.0E-81	3.0E-81	3.0E-81	3.0E-81	2.0E-81	2.0E-81	2.0E-81 AW	2.0E-81	
Expression Signal	3.58	0.98	2.26	2.26	1.33	1.11	0.57	3.59	3.43	3.43	6.1	1.53	17.0	0.71	3.2	3.2	3.63	3.63	1.53	1.53	4.82	12.36	12.36	1.23	5.83	5.83	2.87	2.97	0.71	9.0	
ORF SEQ ID NO:	28580	23013	29509	29510	29733	32834	33058	34016			34980	35861	35935	35936	37086	37087	30731	30732	31030	31031	31004	26658	26659	27832	28400	28401	28254	28255	29144	33678	
Exan SEQ ID NO:	15931	16372	16881	l	17098	19863	19982	20879	21135	21135	21814	22649	22718	22718	23807	23807	25280	25280	24831	24831	24911	13993	13933	15093	15755	15755	15605	15605	16507	20552	
Probe SEQ ID NO:	3168	3619	4139	4139	4360	2212	7289	8185	8443	8443	9128	10001	10070	10070	11140	11140	11928	11928	12463	12463	12597	1244	1244	2371	2989	2989	2837	2837	3755	7857	

Page 380 of 536 Table 4 Single Exon Probes Expressed

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Top Hit Descriptor	23.87 Urmana - di	EST372729 MAGE resembles ANOE U	Zk45f09.r1 Seares_pregnatives, MACF notific suplens CUNA Zk45f09.r1 Seares_pregnant_uterus_NbHPU Horm saplens CDNA clone IMAGE:485825 6' similar to
 | 2002/37888E1 NILL MCC 92 U. | Homo geniens phombodication

 | Homo sapiens phosphodiesterans 10, calmodulin-dependent (70kD) (PDE1C), mRNA
 | Homo sapiens caveolin-1/-2 locus, Contig1, D7S522, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and | 4) Homo conform mail most of films in the state of the st | SOURCESTAIN MOS ES US. | 60164505151 NH MPC 64 U | 601343180F1 NIH MGC 53 Home septems cDNA close IMAGE:3830228 6 | ac14d06.s1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA done IMAGE:866427 3' similar to SW:/YB36_YEAST P38126 HYPOTHETICAL 60.5 KD PROTEIN IN RPS101-RPS13 INTERGENIC
 | 301577338E1 NIH MCC 6 H | 301577339F1 NIT JMGC 9 TOURD SEPTEND CLINA GONE IMAGE:3838280 G | CM3-NN0059-14040n-147-47 Milogo II | Amo sapiens addin-like protein (CLID) with A
 | MR0-CT0008-260599-019 CT0008 Home confee 2014 | JRO-CT0006-250599-019 CT0008 Homo saniens - DNA | ST372729 MAGE resequences. MAGF Home septems CDNA | 101867714F1 NIH MGC 17 Homo saplens cDNA clone IMAGE 24117486 F | tomo sapiens phorbolin (similar to apoliponotein R mRNA editing apolipon 2000) | Homo saplens HSPC288 mRNA, partial cds
 |
| Top Hit
Database
Source | EST DIMANI | EST HUMAN | EST HIMAN | FST HIMAN | LV | Į. | Ę | HST HIMAN | Į. | 12
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| | W26539 1 | AW960658.1 | AA040370.1 | BE047998.1 | U87928.1 | | | AA255569.1 | U62351.1 | U52351.1
 | BF674641.1 | 11420985

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| Most Similar
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Value | 1.0E-81 | 1.0E-81 | 1.0E-81 | 1.05-81 | 1.0E-81 | 1.0E-81 | 1.0E-81 | 1.0E-81 | 1.0E-81 | 1.0E-81
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 | 1.0E-81 / | 1.0E-81 | 1.0E-81 | 1.0E-81 | 1.0E-81 | 8.0E-82
 |
| Expression
Signal | 1.13 | 1.07 | 3.56 | 6.99 | 4.69 | 4.1 | 4.1 | 0.85 | 3.47 | 3.47
 | 1.81 | 0.59

 | 0.59
 | 0.87 | 8.45 | 5.09 | 6.09 | 5.08 | 1.59
 | 3.27 | 3.27 | 1.59 | 2.9
 | 1.97 | 1.97 | 1.72 | 1.99 | 4.13 | 3.6
 |
| ORF SEQ
ID NO: | | | | | | | 30705 | 30922 | 31273 | 31274
 | 31796 | 32218

 | 32219
 | 32416 | 33444 | 35523 | 35624 | 35715 | 35863
 | 35865 | 35866 | 36269 | 36787
 | 36966 | 36967 | 29037 | 37421 | 31093 | 26462
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 | 19401 | 20333 | 22328 | 22328 | 22519 | 22651
 | 22653 | 22653 | 23051 | 23540
 | 23700 | 23700 | 16397 | 24108 | 24622 | 12839
 |
| Probe
SEQ ID
NO: | 1402 | 3644 | 4479 | 4600 | 5157 | 6269 | 5269 | 5415 | 5568 | 5568
 | 6054 | 6453

 | 6453
 | 6639 | 7669 | 9676 | 9876 | 9869 | 10003
 | 10005 | 10005 | 10405 | 10860
 | 11029 | 11029 | 11240 | 11507 | 12132 | 12
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| | Exon ORF SEQ Expression (Top) Hit Top Hit Acesson No. Signal BLASTE No. Source | Exon ORF SEQ ID ID NO: Expression Signal (Top) Hit Top Hit Acesslon Top Hit Acesslon Top Hit Acesslon Top Hit Acesslon Top Hit Acesslon Top Hit Acesslon Top Hit Acesslon Database Source NO: Signal BLAST E No. Source Source 14149 26829 1.13 1.0E-81 W/26539 EST LIMANA | Exch
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FST377729 MAGE Teaching and | Exon
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NO:
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Signal
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Single Exon Probes Expressed in Brain

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Top Hit Descriptor	Homo sapiens HSPC288 mRNA, partial ods	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Homo sapiens mRNA for KIAA1327 protein, partial cds	Homo saplens glutathlone peroxidase 5 (epididymal androgen-related protein) (GPX5), transcript variant 2,	MKINA	Homo sapiens hypothedical protein FLJ20461 (FLJ20461), mRNA	601488531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	AU144050 HEMBA1 Homo sapiens cDNA clone HEMBA1000752 3'	ae68e04.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:969342 3'	nf69er1.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:925196 3'	Homo saplens alpha-tubulin isoform 1 mRNA, complete cds	QV2-HT0540-120900-362408 HT0540 Hamo sapiens cDNA	QV2-HT0540-120900-362-f08 HT0540 Homo sapiens cDNA	Human von Willebrand factor gene, exon 9	wp75e09.x1 NCI_CGAP_Brn25 Homo eaplene cDNA clone IMAGE:2467624 3' similar to TR:075276 075276 PKD1:	Homo saplens presentiin-1 gene, exons 1 and 2	Homo sapiens emyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	RC2-BN0120-010400-013-f02 BN0120 Homo septens cDNA	Homo saplens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo sabiens emyloid beta (A4) precursor protein (probase nexin-)]. Alzheimer disease) (APP), mRNA	ai23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343648 3'	RC6-PT0001-190100-021-B02 PT0001 Homo saplens cDNA	Homo sapiens chromosome 21 segment HS21 C085	RC1-BN0005-260700-018-904 BN0005 Homo saplens cDNA	Homo sapiens adenylate cyclase activating polypeptide 1 (pituitary) receptor type I (ADCYAP1R1) mRNA	Homo sapiens neurotrophic tyrosine kinase, receptor, type 2 (NTRK2) mRNA
Top Hit Database Source	LN	LΝ	Ą	Ą	NT		N.			EST_HUMAN	EST_HUMAN	EST_HUMAN	Ę	EST_HUMAN	EST_HUMAN	N	EST HUMAN	Ę		T_HUMAN			T_HUMAN	Г		EST_HUMAN		
Top Hit Acessian No.	12 AF161406.1	8.0E-82 U08988.1	8.0E-82 U08988.1	2 U08988.1	8.0E-82 AB037748.1			8923432 N	2 BF035327.1	AU144050.1	7.0E-82 AA663747.1	5.0E-82 AA515512.1	4.0E-82 AF081484.1	4.0E-82 BF351691.1	BF351691.1	4.0E-82 M25833.1	4.0E-82 Al937300.1	4.0E-82 AF029701.2	4502166 NT	3E005705.1	3.0E-82 5174702 NT	4502166 NT	3.0E-82 AA725848.1	3.0E-82 AW875073.1	3.0E-82 AL163285.2	3.0E-82 BE813232.1	4501922 NT	5453811 NT
Most Similar (Top) Hit BLAST E Value	8.0E-82	8.0E-82	8.0E-82	8.0E-82	8.0E-82	200	8.0E-82	8.0E-82	7.0E-82	7.0E-82	7.05-82	5.0E-82	4.0E-82	4.0E-82	4.0E-82	4.0E-82	4.0E-82	4.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82
Expression Signal	2.45	1.1	2.83	0.84	1.42	,	1.43	6.0	1.7	1.2	1.71	0.71	49.82	0.8	8.0	0.65	11.61	5.05	21.65	3.11	5.7	10.65	18.58	1.25	3.44	1.91	1.53	2.52
ORF SEQ ID NO:	25452	25704	26227	26306	26907		27089	CSCSZ		28216	37682	29473	27103	30915	30916	31381	37633		25721	26110	26203	26290		26758	26881	27341	27459	
Exan SEQ ID NO:	12839		13567		14221		14400	16960	14180	15474	24350	16847	14412	18208	18208	18466	24310		13079	13462	13542	13620	13799	14082	14197	14631	14735	16028
Probe SEQ ID NO:	104	257	795	867	1474		1634	4219	· 1433	2769	11759	4104	1666	5409	5409	5671	11716	12374	271	687	770	850	1039	1333	1450	1894	2000	3266

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Top Hit Descriptor	zn93b04.r1 Stratagene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:565711 ff' similar to	SW.PAGT_BOVIN Q07837 POLYPEPTIDE N-ACETYLGALACTOSAMINY TRANSEERASE	Homo sapiens ankyrin-like with transmembrane domains 1 (ANKTM1) mRNA	Homo saplens contactin 6 (CNTNB) mRNA	Homo sapiens contactin 6 (CNTNR) mRNA	Homo sepiens mRNA for KIAA1077 protein partial cds	Homo sapiens mRNA for KIA41077 protein natial cab	Homo sapiens mRN4 for KL4.0000 postata adv	Homo sapiens mRNA for KIAAngoo natein partein ada	DKFZp434M117 r1 434 (swnown: hises) Home contract chiving class chiving classifications	Homo saniens DNA for emildid programmer profess county cities and county of the county	Homo saplens alutamate receipts Innotants Points 4 (Celiza) - Data	Homo sepiens mRNA for KIAA4nog arctein and Ad	Homo saplens mRNA for KIAA1096 protein, partial ords	Homo sapiens wbscrt (WBSCR1) and wbscr6 (WBSCR6) genes, complete cds, alternatively spliced and	Homo saplens himor nacmala faster according and a complete cas	Homo sablens timos negrosts (social receptor superillaminy, member 5 (TNFKSF8) mRNA	Homo sepiens mRNA for KIAA0727 protein pertein of a	Homo saplens FAM4A1 splice variant a (FAM4A1) mPNA	tm21g05.x1 Soares NFL T GBC S1 Homo sapiens cDNA chas link ACE 21 #2272 91	Homo sapiens hypothetical protein FLJ20128 (FLJ20128), mRNA	Homo sapiens alt (Drosophila) homolog 3 (SLIT3), mRNA	Homo sapiens microrchidia (mouse) homolog (MORC), mRNA	Homo saplens microrchidia (mouse) homolog (MORC), mRNA	Human endogenous retrovirus-K, LTR U5 and agg gene	Human endogenous retrovirus-K. LTR U5 and gap gene	Homo saplens leucy/cvstfnvl emhonentidase (1 NDED) mBNA	Homo sapiens leucy/cystiny aminopentidase (I NPED) mPNA	Homo sapiens CAGF9 mRNA, partial cds	Homo sepiens CAGE9 mRNA, partial cds	Homo sapiens EGF-like repeats and discoidin Flike domains 3 (EDII 3) mRNA	2b31d10.s1 Sceres_parethyroid_tumor_NbHPA Homo saplens cDNA clone IMAGE:305203 3'
Top Hit Database Source		EST HUMAN	LN	N	Ρ	LN LN	F	LN	N.	EST HUMAN		1Z	Į.	Z.		Ę	Į.	Į	ĮN.	EST HUMAN	Z	ΤN	F	F	NT	N	5	5	N	N	5	EST_HUMAN
Top Hit Acession No.		-82/AA135979.1	11425206 N	11432889 NT	11432889 NT	-82 AB029000.1	-82 AB029000.1	-82 AB023216.1	-82 AB023216.1	82 AL046390.1	82 D87675.1	4504118 NT	82 AB029019.1	82 AB029019.1	82 AF045555 1	4507580 NT	4507580 NT	82 AB018270.1		Γ	2.0E-82 8923130 NT	11321570 NT	7657340 NT	7657340 NT	П		11417191 NT	11417191 NT			5031660	
Most Similar (Top) Hit BLASTE Value		3.0E-82	3.05-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82 Y	2.0E-82 Y	2.0E-82	2.0E-82 11	2.0E-82 U80736.1	2.0E-82 U80736.1	2.0E-82	2.0E-82 N94950.1
Expression Signal		0.98	3.14	0.88	0.88	3.23	3.23	2.55	2.55	1.21	1.25	1.17	1.01	1.01	2.85	1.46	1,46	5.65	4.73	0.91	0.85	2.18	0.45	0.45	1.84	1.84	1.27	1.27	4.45	4.45	19.1	1.58
ORF SEQ ID NO:								25990	25991	27121	29210	29575	29890	29891	30172	30352	30353	30874	31827		33482	34035	34397	34398	35870	35871	37187	37188	37233	37234	37870	
SEQ ID NO:	5	1/283	20746	_]	- 1	ı	١			ı	16578		17256	17256	17547	17742	17742	18184	18861	25426	20369	20898	21280	21260	22857	22657	23900	2300	23940	23940	24341	24508
Probe SEQ ID NO:	200	5	8002	8464	8454	9724	9724	983	583	1681	3827	4208	4521	4521	4816	5021	5021	5384	6082	7581	7705	8204	8268	8268	10009	1000	11237	11237	11279	11279	1750	11957

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	Top Hit Descriptor	zi01g09.r1 Scares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:429568 5	Homo sapiens SRY (sex determining region Y)-box 10 (SOX10), mRNA	Homo sapiens melanoma differentiation associated protein-5 (MDA5), mRNA	601510859F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912207 5	RC4-BT0310-110300-015-f10 BT0310 Homo saplens cDNA	Homo sapiens mRNA for KIAA0538 protein, partial cds	Homo sapiens mRNA for KIAA1417 protein, partial cds	Homo sapiens mRNA for KIAA0662 protein, partial cds	UI-H-BW1-aca-f-03-0-UI.s1 NCL_CGAP_Sub7 Homo sapiens cDNA clone IMAGE:3084053 3'	Homo sapiens chromosome 21 segment HS21C009	602150403F1 NIH_MGC_81 Hamo saplens cDNA clone IMAGE:4291561 5'	301117160F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3357734 5'	601273346F1 NIH MGC_20 Homo sapiens cDNA clone IMAGE:3814382 5'	za48f12.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295823 3'	QV4-LT0016-271299-068-h11 LT0016 Homo sapiens cDNA	no12h01.s1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100497 3' similar to contains Alu repetitive element:	7p37a07.x1·NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:3647893 3' similar to TR:Q9Y3I6 Q9Y3I6	DJ207H1.1;	Homo sapiens KIAA0100 gene product (KIAA0100), mRNA	Homo sapiens transcription factor CA150 (CA150) mRNA	Homo sapiens transcription factor CA150 (CA150) mRNA	Human platelet Glycoprotein IIb (GPIIb) gene, exons 2-29	hf31h03.x1 Soares_NRT_GBC_S1 Homo saplens cDNA clone IMAGE::2933525 3' similar to SW:YBEB HAEIN P44471 HYPOTHETICAL PROTEIN HI0034. ;	QV4-ST0234-181199-037-105 ST0234 Homo saplens cDNA	Homo saplens chromosome 21 unknown mRNA	Homo sapiens VAMP (vesicle-associated membrane protein)-associated protein A (33kD) (VAPA) mRNA,	and translated products	Homo sapiens hyperion gene, exons 1-50	Homo saplens met proto-oncogene (hepatocyte growth factor receptor) (MET), mRNA	Homo sapiens myomesin (M-protein) 2 (165kD) (MYOM2), mRNA	Homo sapiens pre-mRNA splicing factor similar to S. cerevisiae Prp18 (PRP18), mRNA	11 Date Date
	Top Hit Database Source	T_HUMAN			EST_HUMAN	T_HUMAN	TN		LN	T_HUMAN			HUMAN		EST_HUMAN	EST_HUMAN	HST HUMAN	T	T_HUMAN				LN	EST HUMAN	Γ	Г			NT				
*	Top Hit Acession No.	2.0E-82 AA011278.1	11418097 NT	11545921 NT	1.0E-82 BE885106.1	1.0E-82 BE064386.1	1.0E-82 AB011110.2	1.0E-82 AB037838.1	1.0E-82 AB014562.1	1.0E-82 BF515938.1	1.0E-82 AL163209.2	9.0E-83 BF672220.1	9.0E-83 BE253347.1	8.0E-83 BE383973.1	8.0E-83 N66951.1	7.0E-83 AW385529.1	7 0F-83 AA584655 1		7.0E-83 BF221813.1	11426657 NT	5729753 NT	5729753 NT	6.0E-83 M33320.1	6.0E-83 AW573088.1	6.0E-83 AW816405.1	6.0E-83 AF231919.1		4507866 NT	6.0E-83 AJ010770.1	11422024 NT	4505314 NT	11430647 NT	214 -1 -1 -1 -1
	Most Similar (Top) Hit BLAST E Value	2.0E-82	2.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	1.0E-82	9.0E-83	9.0E-83	8.0E-83	8.0E-83	7.05-83	7.05-83		7.0E-83	7.0E-83	7.0E-83	7.0E-83	6.0E-83	6.0 = -83	6.0E-83	6.0E-83		6.0E-83				6.0E-83	100
	Expression Signal	3.47	1.95	1.14	0.77	3.1	1.28	1.13	0.59	1.17	2.34	4.51	0.53	3.33	5.63	26.0	88		89.9	0.58	1.1	4.1	1.98		0.71	1.08		2.02	1.52	2.27	2.85	2.34	,,,,
	ORF SEQ ID NO:			25985		26679	26680	34678	35390		36576	34449	36039	26816	27115	26759				31702	37634	37635	25826	27224	28432	-		30641	31674	33160	35413	35517	0,100
	Exan SEQ ID NO:	24844	25029	13358	13938	L	ı	21533	1			ı	ı	l	15523	14084	15835		17497	L	24311		13179	14520				18019	18716	20079	1		ı
	Probe SEQ ID NO:	12485	12775	578	1186	1263	1264	8841	9553	10145	10646	8615	10174	1392	1676	1335	2868		4765	2969	11717	11717	394	1770	3017	3046		5211	5633	7401	9675	6996	3

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Olingie Exort Probes Expressed in Brain	Top Hit Descriptor	ab14e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:840810 3' similar to contains	Home septent spectrum of the septent	garles, complete cas	numen succinate denydrogenase fron-protein subunit (sdhB) gene, exon 5	Nowel himen and manufactured and regulatory subunit (SUG2) mRNA, complete cds	Homo seriens demostisment	Home capture of the control of the control of the capture of the c	Homo sapiens catalana (CAT) - ENIA	Homo sepiens E.GE.IIIA paraceta and discrete in the control of the	Home capters ECE III.	Honno septents E.S.Ftike repetits and discodin Filite domains 3 (EDIL3), mRNA Honno septens mannosidasse, beta A, lysocomal (MANBA) gene, and ublquitin-conjugating enzyme E2D 3	(VOLLAND) genes, complete cds	np87c07.s1 NCI_CGAP_Thy1 Homo sapiens cDNA similar to endogenous retrovins ERV9	repetitive element;	of64g05.s1 Soares_tests_NHT Homo sapiens cDNa clone IMAGE:1765682.3/	Q92614 MYELOBLAST KIAA0216.;	0084g05.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1621592.3' similar to TR:Q92614	2848/12.51 Soares fatal liver sploon 4NET 8 U.S.	RC6-ET0046-280600-013-H12-ET0048 Home capiens cDNA clone IMAGE:285823 31	Homo sapiens sai (Drosophila)-like 1 (SAI 11) mRNA	Homo sablens chromosome 21 serment LEST COO	Homo saplens hematopoletic progenific cell antinen CD24 accession 2014	Homo septens enkirtin reneal control in a result of the control of	Homo septiens ankwin repeat-containing protein ASD-2 (LOCO16/8), mRNA	lumen cardiocambhanic matica	Homo senties membrane state: CH4 / CH4	0150748751 NIH MGC 74 Home Collins Col	Homo saplens F-box protein Fbl3b/FBl 38) mBNA modulade: 3909068 6'	Homo saplens F-box protein Fbl3b (FBL3B) mRNA, partial cds
וטום באטוו גיוס	Top Hit Database Source	EST HUMAN	- L	į	NT.	FN	₽Z	Į.	LZ	Z	Į.	L L	EST HUMAN		Т	Т	EST_HUMAN	EST HUMAN	Т	Т		LN						T HUMAN		TN T
בֿל ו	Top Hit Acession No.	-83 AA486105.1	-83 AF240788 1	-83 U17883 1	5.0E-83 AF006305 1	5.0E-83 AL 133207.2	4885190 NT	4557013 NT	4557013 NT	5031660 NT	5031660 NT	83 AF224669 1			T		B3 AA993492.1			2.0E-83 BE828694.1	834		2.0E-83 AF202879.1	TV06398 NT	7706398 NT	08879.1	428081			П
	Most Similar (Top) Hit BLAST E Value	6.0E-83	6.0E-83	5.0E-83	5.0E-83	5.0E-83	5.0E-83	5.0E-83	5.0E-83	5.0E-83	5.0E-83	4.0E-83	3.0E-83/	3 OF 83	3 0F-83 /		2.0E-83	2.0E-83	2.0E-83 N	. 2.0E-83 B	2.0E-83	2.0E-83 A	2.0E-83 A	2.0E-83	2.0E-83	2.0E-83 U06679.1	2.0E-83	2.0E-83 BE885401.1	2.0E-83 AF129533.1	2.0E-83 AI
	Expression Signal	2.53	4.27	203	1.55	1.18	0.77	11.53	11.53	1.07	1.07	1.72	4.9	1.33	0.62		1.86	1.86	4.07	1.1	1.89	0.7	4.11	6.14	6.14	0.0	0.85	1.31	1.12	6.36
	ORF SEQ ID NO:						29275	30350	30351	30428	30429	26039					27240	27241	27365	28268	-		29668	29968	29969	30620	31468	31603	32423	33095
	Exan SEQ ID NO:	24117	24472	13692	15526		- 1	İ	- 1	- 1	17812	13404	13742	15485	19250	7,697	14032	14632	14655	15624	16025	16508	17041	1/338	17339	17997	18547	18662	19409	20017
	Probe SEQ ID NO:	11517	11908	926	2043	3629	3886	60 20 20 20 20 20 20 20 20 20 20 20 20 20	5020	508	5093	625	977	2780	6483	1702	78/	1792	1918	2826	2000	3/20	4302	\$ E	4604	5189	6765	5875	6647	7335
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	THE PARTY OF THE P
Top Hit Descriptor 801822090F1 NIH_MGC_75 Home saplens cDNA clone IMAGE:4042318 5' Home saplens mRNA for brain ryanddine receptor, complete cds Home saplens mRNA for brain ryanddine receptor, complete cds Home saplens mRNA for brain ryanddine receptor, complete cds Home saplens mRNA for brain ryanddine receptor, complete cds Home saplens protein kinase CK2 catalytic subunit alpha gene, exen 1 Home saplens protein kinase CK2 catalytic subunit alpha gene, exen 1 Home saplens phosphorylase kinase, gamma 1 (muscle) (PHKG1) mRNA, 3' end Home saplens phosphorylase kinase, gamma 1 (muscle) (PHKG1) mRNA, 3' end Human neural cell adhesion molecule (N-CAM) secreted isoform mRNA, 3' end AU117859 HEMBA1 Home saplens cDNA clone HEMBA1001910 5' ULHF-BNO-and-h-07-0-UI.r1 NIH_MGC_50 Home saplens cDNA clone IMAGE:3081852 5' Home saplens KIAA0985 protein (KIAA0985), mRNA DKFZp547J135_r1 547 (synonym: hfbr1) Home saplens cDNA dene DKFZp547J135 5' Home saplens gene for AF-6, complete cds Home saplens gene for AF-6, complete cds	hydratase (trifunctional protein), beta subunit (HADHB) mRNA Homo sapiens hydroxyacyl-Coenzyme A dehydrogense/3-ketoacyl-Coenzyme A thiolase/enoyl-Coenzyme A hydratase (trifunctional protein), beta subunit (HADHB) mRNA hydratase (trifunctional protein), beta subunit (HADHB) mRNA, complete cds Homo sapiens lipopolysaccharide-binding protein (LBP) mRNA, complete cds Homo sapiens lipopolysaccharide-binding protein (LBP) mRNA, complete cds Homo sapiens loal recognition molecule Caspr2 (KIAA086B), mRNA, partial cds Homo sapiens predic cortactin-binding protein CBP90 mRNA, partial cds H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens gene for mitochondrial dodecencyl-CoA delta-tsomerase, exon 3 H. sapiens cDNA clone IMAGE:3958853 5° H. sapiens cDNA for 19-20600-011-g05 FN0119 Homo sapiens cDNA RC2-FN0119-200600-011-g05 FN0119 Homo sapiens cDNA ae886a03.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:971020 3° E. Sapiens delta-construction for the sapiens cDNA clone IMAGE:971020 3° E. Sapiens delta-construction for the sapiens cDNA clone IMAGE:971020 3° E. Sapiens delta-construction for the sapiens cDNA clone IMAGE:971020 3° E. Sapiens delta-construction for the sapiens cDNA clone IMAGE:971020 3° E. Sapiens delta-construction for the sapiens cDNA clon
Top Hit Database Source Source EST HUMAN NT NT NT NT NT NT NT NT NT NT EST HUMAN NT NT EST HUMAN NT EST HUMAN NT NT NT NT NT NT NT NT NT NT NT NT NT	NT NT NT NT NT NT EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN
Top Hit Acession No. No. No. BF105097.1 ES AB001025.1 NT AB001025.1 NT AB001025.1 NT AF011920.1 NT AF011920.1 NT AF011920.1 NT AF011920.1 ES A453881 NT M22094.1 NT M22094.1 NT AL134448 NT AL134448 NT AL134452.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES AW501399.1 ES	4504326 NT 4504326 NT AF105087.1 NT AF105087.1 NT Z25822.1 NT Z25822.1 NT Z25822.1 NT Z25822.1 NT Z25822.1 ES BE901209.1 ES BE838864.1 ES BE838864.1 ES AA776574.1 ES AA776574.1 ES
	1.0E-83 1.0E-8
Expression Signal 0.044 0.078 0.748 0.048 4.01 1.39 0.77 0.77 0.77 0.77 0.77 0.77 0.77 0.7	2.18 2.18 3.99 3.39 3.38 3.38 3.38 3.38 3.38 3.3
<u>8</u> 0	20814 20874 20874 20874 20873 20874 20873 20874 20167 20167 20167 20167 20167 20167 20167
SEQ ID NO: 20367 20438 20438 20907 22147 22147 22758 22582 22582 22582 23537 23438 23527 2	14137 14189 14189 15842 16860 16860 16860 16860 16860 16828 14021 14021 15117 17892
Probe NO: 7704 7742 7742 7744 7744 7744 7744 7744	1390 1390 1442 1442 1442 1442 1442 1442 1442 144

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P0667 al47g03.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1460500 3' similar to gb:M14338 243997.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:632100 6' similar to TR:G483915 G483915 RETROTRANSPOSABLE L1 ELEMENT LREZ FROM CHROMOSOME 10.; Homo saplens acetyl LDL receptor; SREC≃scavenger receptor expressed by endothelial cells (SREC), Homo saplens acetyl LDL receptor; SREC=scavenger receptor expressed by endothelial cells (SREC), Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, wa76c04.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2302086 3' cimilar to Homo sepiens X-linked Juvenile retinoschists precureor protein (XLRS1) mRNA, complete ods Homo sapiens regulatory factor X, 3 (influences HLA class II expression) (RFX3), mRNA Homo saplens discs, large (Drosophila) homolog 2 (chapsyn-110) (DLG2) mRNA Homo seplens myosah light chain kinase isoform 2 (MLCK) mRNA, complete cds Homo saplens protein tyrosine phosphatase, receptor type, G (PTPRG), mRNA Homo saplens protein tyrosine phosphatase, receptor type, G (PTPRG), mRNA Homo saplens histone deacetylase 3 (HDAC3) gene, complete cds Homo saplens KIAA0783 gene product (KIAA0783), mRNA Homo sepiens discs, large (Drosophila) homolog 2 (chapsyn-110) (DLG2) mRNA Homo sapiens mRNA for KIAA1130 protein, partial cds PM0-LT0019-190600-004-F02 LT0019 Homo saplens cDNA Homo saplens pro-mRNA splicing factor (PRP18) mRNA, complets cds PM4-FT0054-160600-004-e10 FT0054 Homo seplens cDNA VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN); Top Hit Descriptor Novel human mRNA containing Zinc finger C2H2 type domains SW:NRDC_HUMAN 043847 NARDILYSIN PRECURSOR Homo sapiens tropomodulin 2 (neuronal) (TMOD2), mRNA Homo sapiens Bach1 protein homolog mRNA, partial cds Homo sapiens mRNA for KIAA1131 protein, partial cds Homo sapiens mRNA for KIAA1131 protein, partial cds Homo sapiens pericentriolar material 1 (PCM1) mRNA Homo sapiens chromosome 3 subtelomeric region EST96094 Tests I Homo saplens cDNA 5' end Single Exon Probes Expressed in Brain complete cds) mRNA mRNA EST_HUMAN EST_HUMAN EST_HUMAN Top Hit Database HUMAN EST_HUMAN EST_HUMAN Source EST. 11428740 NT 957.1 NT Ę 11426718 NT 뉟 눌 11386168 NT 눋 뉟 5453855 NT 11426718 Top Hit Acesslon 11433550 11421326 4557528 4557526 6.0E-84 AA897339.1 AB032957.1 6.0E-84 AF038391.1 6.0E-84 BE770199.1 5.0E-84 AA382811.1 5.0E-84 AF109718.1 5.0E-84 AB032957.1 5.0E-84 AB032957.1 4.0E-84 AF059650.1 4.0E-84 1142 4.0E-84 455 5.0E-84 AA167678.1 4.0E-84 AI685321.1 AF069601.2 3.0E-84 AF028200.1 3.0E-84|AB026898.1 3.0E-84 AL096880.1 3.0E-84 AF014459.1 6.0E-84 6.0E-84 6.0E-84 (Top) Hit BLAST E 4.0E-84 Most Simila 8.0E-84 5.0E-84 4.0E-84 4.0E-84 Value 1.87 Ģ 40. 0.97 2.37 0.59 3.17 2.19 14.38 0.71 1 1.36 1.88 1.07 Expression 5.2 1.21 4.51 Signal ORF SEQ 30943 31282 33134 33367 33789 31283 30242 31168 26121 37438 37570 37571 37738 31925 34647 33319 25752 27401 28977 29121 ÖΝΩ SEQ ID 18230 13472 18371 20053 20259 18796 24249 18371 24133 24249 18952 13112 16333 20217 23517 14688 16483 14736 1827 21501 ÿ 1577 Probe SEQ ID 5431 7373 7591 7972 6015 5574 5574 1389 4897 5475 3013 1633 8809 697 11652 11652 5475 6175 7547 10835 1953 3578 3731 289

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Top Hit Descriptor	wu20d05x1 Soeres, Dieckgreefe, colon, NHCD Homo sapiens cDNA clone IMAGE:2520585 3' similar to gb:L050s3 60S RIBOSOMAL PROTEIN L18A (HUMAN);	CM1-BT0795-190600-272-b08 BT0795 Homo sapiens cDNA	CM1-BT0795-190600-272-b08 BT0795 Homo sapiens cDNA	Homo sapiens myelin transcription (actor 1-like (MYT1-l) mRNA, complete cds	H. sapiens DNA for endogenous retroviral like element	ULH-BI4-acl-a-02-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3084963 3'	UI-H-BI4-ad-a-02-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3084963 3'	yr56e11.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:209324 3'	qm87c09.x1 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1895728 3'	Homo saplens chromosome 21 segment HS21C004	Homo saplens chromosome 21 segment HS21C004	AU120280 HEMBB1 Homo sapiens cDNA clone HEMBB1000339 5	ym49e11.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:51383 5' similar to SP.APOH_RAT P26644 BETA-2-GLYCOPROTEIN I	nae30e02x1 Lupski, sympathetic, trunk Homo sapiens cDNA clone IMAGE:4090251 3' similar to TR:Q9UGS3 Q9UGS3 DJ756G23.1;	nae30a02x1 Lupski, sympathetic, trunk Homo sapiens cDNA clone IMAGE:40902513' similar to TR:09UGS3 Q9UGS3 DJ756G23.1;	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta polypeptide (YWHAZ) mRNA	Homo sapiens complement component 5 (C5), mRNA	am85b11.s1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:1629885 3'	601308006F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3626257 5	Homo sapiens pericentriolar material 1 (PCM1), mRNA	nw12e06.s1 NCI_CGAP_SS1 Homo sapiens cDNA clone IMAGE:1239106 3'	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	DKFZp434N0323_r1 434 (synanym: htes3) Hamo sapiens cDNA clone DKFZp434N0323 5'	DKFZp434N0323_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434N0323 5	Homo sapiens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Homo saplens catenin (cadherin-associated protein), alpha 2 (CTNNA2), mRNA	Homo sapiens speckle-type POZ protein (SPOP), mRNA	
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	卜	F	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT TN	Ę	LN	EST_HUMAN	EST_HUMAN	TN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	NT	TN	NT	
Top Hit Acession No.	Al983801.1	BE695397.1	BE695397.1	AF036943.1	X89211.1	BF511575.1	BF511575.1	H63370.1	AI298674.1	AL163204.2	AL163204.2	AU120280.1	H22841.1	BF448000.1	BF448000.1	AF114488.1	4507952 NT	11427631 NT		BE392137.1	11427197 NT	AA720851.1	AJ229041.1	AL043314.2	AL043314.2	AJ229041.1	7656998 NT	11434422 NT	
Most Similar (Top) Hit BLAST E Value	3.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84	2.0E-84		2.0E-84	2.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84 /	1.0E-84 /	1.0E-84	1.0E-84	
Expression Signal	3.55	6.94	6.94	9.31	0.77	0.92	0.92	0.75	1.35	0.49	0.49	0.81	0.61	<u></u> е	8	1.89	20.64	٦	3.17	1.92	1.13	2.46	5.01	3.53	3.53	2.67	1.15	0.98	
ORF SEQ ID NO:		27563	27564	28362	28378	30952	30953	32310		34118	34119	35094	35476	31100	31101	25748	25953		26685	27508	27686	29123	29752	30017	30018	29752	30483	31551	
Exon SEQ ID NO:	23473	14829	14829	15710	15728	18238	18238	19305	20651	20978	20978	21924	22283	24643	24643	13108	13319	13478	14019	14781	14948	16486	17120	17385	17385	17120	17870	18619	
Probe SEQ ID NO:	10790	2098	2098	2944	2962	6439	5439	6540	7956	8284	8284	9245	9631	12159	12159	304	536	703	1270	2048	2220	3733	4383	4851	4651	4855	5153	5830	

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	1340								I											4	†	ا "	"		1	7		*	C		567
Top Hit Descriptor	uterine water channei≕28 kda erythrocyte integral membrane protein homolog [human, uterus, mRNA, 1340] nti	Novel human gene mapping to chomosome 13	Novel human gene mappling to chomosome 13	Novel human gene mapping to chomosome 13	Homo saplens polymerase (DNA directed), alpha (POLA), mRNA	Homo sapiens NGFI-A binding protein 1 (ERG1 binding protein 1) (NAR1) mBNA	Homo saplens nuclear transport factor 2 (placental protein 15) (Pp15) mRNA	Homo saplens Ca2+-binding protein CABP3 (CABP3) gene, exon 6 and nartial cds	Homo sapiens ubiquitin specific protease 13 (Isopeptidase T-3) (USP13) mRNA	Homo sapiens ublquitin specific protease 13 (Isopentidase T-3) (ISP13) mRNA	Homo sapiens glutamate receptor, ionotrophic, AMPA 4 (GR)441 mRNA	Homo saplens purinergic receptor P2X-like 1 oranga receptor (P2RX13) mbnis	Homo sapiens aconitase 2, mitochondria (ACO2), mRNA	Homo saplens chromosome 21 segment HS21Cong	Homo sapiens nuclear protein Skip mRNA complete cds	Homo sepiens nuclear protein Skin mRNA Apmilia Ada	Homo sablens (eubarth (LDPI) mRNA	Human plasminoden dene, exon 7	Human plasminoden gene exon 7	Homo saplens DKF7p434P311 prolein (DKF7b434D341)DNIA	Homo sepiens chromosome 21 segment HS21Chan	Homo saplens heat shock transcription factor 2 hinding protein (LISESDB)DN A	Homo saplens chromosome 21 segment HS21Ch88	Homo sapiens ribosomal protein L27 mRNA, complete cde	Homo sapiens MSTP030 mRNA, complete cds	Homo sapiens DEAD/H (Asp-Gil-Ala-AspiHis) how notweetide to (DNA holls-of) How	UNITED TO THE COURT OF THE COUR	ronno septens DEADM (ASP-ciu-Ala-Asp/Mis) box polypeptide 10 (RNA helicase) (DDX10), mRNA zt62b01.r1 Soares, testis. NHT Homo saplens cDNA clone IMAGE-72889 6' similar to TB-2-13357eo	G1335769 GAG-POL POLYPROTEIN.;	Homo sapiens chromosome 21 segment HS210084	
Top Hit Database · Source	Į Į	Ϋ́	ΙZ	Į.	Ę	N	N	Z	Ę	FZ	Z	Ę	Ę	ŁN	LN LN	Z	5	Ν	LN FX	-	Z	F	F	Į.	-FA	5	Ŀ		EST_HUMAN	П	
Top Hit Acession No.	S73482.1	AL049784.1	AL049784.1	AL049784.1	R393994 NT	11430846 NT	5031984 NT	AF224511.1	4507848 NT	4507848 NT	11437356 NT	11417812 NT	11418185 NT	AL163209.2			4758669	M33282.1		7657020	AL163280.2	1979	AL163268.2		AF113210.1	11438573 NT	11420672 NIT	2		AL163284.2 N	
Most Similar (Top) Hit BLAST E Value	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84				1.0E-84	1.0E-84	1.0E-84	1.0E-84	9.0E-85	9.0E-85			9.0E-85	9.0E-85	9.0E-85			9.0E-85	7.0E-85 L	7.0E-85	6.0E-85	8 HO			5.0E-85 A	
Expression Signal	1.41	1.68	1.66	2:32	1.26	2.42	3.05	0.53	3.05	3.05	1.08	2.34	3.2	1.06	2.39	2.39	0.95	1.23	1.23	3.6	96'0	0.98	1.12	1.45	4.32	3.35	3.35	2	1.29	1.49	
ORF SEQ ID NO:						33229					36368		31096		26469	26470	26783	27004	27005	27108	29591	30177	30208	26530		37300	37301		37690	27792	
SEQ ID	1		\Box	_[_		_[_[\perp				13810	13810	14108		14319	14415	16966	17555	17585	13871	24239	23998	23998		24357	15056	
Probe SEQ ID NO:	6097	6781	6781	7007	7369	7501	9435	9670	0696	0696 6	10496	12046	12151	946	1051	1051	1360	1572	1572	1670	4225	4824	4856	1114	11642	11392	11392		11766	2332	

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	complete cds			гута Е2D 3	complete cds												4		10621 protein								74	Ü		E	67
Top Hit Descriptor	Homo saplens T-type calcium channel alpha1 subunit Alpha11-a isoform (CACNA11) mRNA, complete cds	601458646F1 NIH_MGC_66 Hamo sapiens cDNA clane IMAGE:3862402 5'	601458646F1 NIH_MGC_66 Hamo sapiens cDNA clone IMAGE:3862402 5'	Homo sepiens mannosidase, beta A, lysosomal (MANBA) gene, and ubiquitin-conjugating enzyme E2D 3 (UBE2D3) genes, complete cds	Homo sapiens T-type calcium channel alpha1 subunit Alpha11-e Isoform (CACNA11) mRNA, complete cds	602084730F1 NIH_MGC_83 Hamo sapiens cDNA clone IMAGE:4249087 5"	602084730F1 NIH_MGC_83 Hamo sapiens cDNA clone IMAGE:4249087 51	RC1-BT0623-120200-011-c07 BT0623 Homo saplens cDNA	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 6	ye53g09.r1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:121504 5	601189704F2 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3533616 5	Homo sapiens F-box only protein 24 (FBXO24), mRNA	Homo sapiens F-box only protein 24 (FBXO24), mRNA	Homo sapiens lacrimal proline rich protein (LPRP), mRNA	Homo sapiens met proto-oncogene (hepatocyte growth factor receptor) (MET), mRNA	Homo sepiens KIAA0793 gene product (KIAA0793), mRNA	Homo saplens KIAA0793 gene product (KIAA0793), mRNA	Homo saplens mRNA for dynein heavy chain (DNAH9 gene)	Homo sepiens GTPase regulator associated with the focal adhesion kinase pp125(FAK); KIAA0621 protein ^t (KIAA0621), mRNA	Homo saplens DENN mRNA, complete cds	Homo sapiens CGI-81 protein (LOC51108), mRNA	Homo sepiens phospholipase C, epsilon (PLCE), mRNA	Homo sapiens small nuclear ribonucleoprotein polypeptide B" (SNRPB2), mRNA	Homo sapiens small nuclear ribonucleoprotein polypeptide B" (SNRPB2), mRNA	Homo saplens phospholipid scramblase mRNA, complete cds	RC1+HT0268-031299-012-f09 HT0268 Homo sapiens cDNA	Homo sepiens EGF-like repeats and discoldin I-like domains 3 (EDIL3), mRNA	Homo sapiens mRNA for KIAA1107 protein, partial cds	Homo sapiens mRNA for KIAA1107 protein, partial cds	Homo saplens Ran GTPase activating protein 1 (RANGAP1), mRNA	
Top Hit Database Source	LN	EST_HUMAN	EST_HUMAN	NT	١	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	NT	TN	LN	Z	NT	FZ	NT		Z	N	۲	N	NT	NT	EST_HUMAN	TN.	TN	NT	NT	
Top Hit Acession No.	AF211189.1	BF035674.1	BF035674.1	AF224669.1	AF211189.1	BF677910.1	BF677910.1	BE079263.1	AF096157.1	T97495.1	BE267189.1	11024695 NT	11024695 NT	11436001 NT	11422024 NT	7662309 NT	7662309 NT	AJ404468.1	11416870 NT	U44953.1	11525829 NT	11430889 NT	11421422 NT	11421422 NT	AF098642.1	BE150392.1	5031660 NT	AB029030.1	AB029030.1	11418177 NT	
Most Similar (Top) Hit BLAST E Value	5.0E-85	5.0E-85	5.0E-85	5.0E-85	5.0E-85	4.0E-85		4.0E-85	3.0E-85	3.0E-85		3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.05-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85	3.0E-85		3.0E-85	
Expression Signal	0.8	1.4	1.4	7	5.28	1.51	1.51	1.3	2.98	3.51	6.53	1.45	1.45	1.07	0.63	5.71	5.71	7.79	0.91	1.89	0.74	3.8	0.96	96.0	0.56	1.88	2.25	1.79	1.79	1.98	
ORF SEQ ID NO:			30852	37005		31797	31798			27215		30194	30195	202777	31737	31783	31784		33055		34235			35258	36242	36659	37403		37743		-
Exen SEQ ID NO:	17136	18166	1	23733	17136	18836	18836	23118		14515	17019	17571	17571	18120	18775	18823	18823	19553	19978	L	21099	21568			23027	23418	24091	24408		24937	
Probe SEQ ID NO:	4399	5364	5364	11063	12743	9509	9509	10472	1276	1773	4280	4841	4841	5316	5994	6043	6043	6853	7295	777.1	8406	8877	9208	9206	10381	10730	11490	11824	11824	12640	

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Table 4
Single Exon Prohes Fynnsson

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		i										lement									#	P II					5		9	-		1	I	JE	5
	Top Hit Descriptor	Home confee attendant of (1) to 1000	Home service: CO 201 - 1 - 1 - 201 -	Home september and prometries (LOCat 340), mrnA	Homo sandens engineering CII (APOCZ) many	Human DNA polymerase bets sons 2000 42 4.6	Homo sabiens similar to ref lateural months.	Human Ku (520/580) subunit mBNA annulus 31/50/50/50/50 POM121 (POM121L1), mRNA	Homo sapiens plasminosen (Pi A) m BNA	Homo sablens realin (REIN) mRNA	Homo sabians chromosome 21 segment US21C064	wid7h08.x1 NCI_CGAP_Kid12 Homo sapiens oDNA clone IMAGE.2388431.3' similar to contains element	woldows v comes NET + OBO P. I	WIND WAR CONTROL OF THE WAR CONTROL OF THE WIND WAS CONTROL OF THE WAR CONTROL OF THE WAR CONTROL OF THE WAR CONTROL OF THE WAS CONTROL OF THE WAR	1601591416F1 NIH MAC 7 LOS PRINCES CON COMPANY CON CONTRACT OF CON	1601462817F1 NIH MGC 67 Home semicas CONA Glone IMAGE:3946818 6	601462817F1 NIH MGC 67 Homo sealons of the live 114 OF 200021 5	601109738F1 NIH MGC 18 Home companion ONA 21 111100 0000000000000000000000000000	246f03 s.1 Sories fetal liver enlaw anich S of U.	245(03.s1 Scares felal liver spleas 1MEIS ST Long	601897003F1 NIH MGC 19 Homo seplens of IN A September 501897003F1 NIH MGC 19 Homo seplens of INA September 50189700000000000000000000000000000000000	601897003F1 NIH MGC 19 Home sealests CDNA clotte 117 OF 1255	Human mRNA for T-cell ovclobilin	qi56a07.x1 NCI CGAP Bm25 Homo sanlans cDNA Alone IMAGE: 1880458 gi	Homo saplens calcineurin binding protein 1 (KIAA0330) mBNA	Homo saplens calcineurin binding protein 1 (KIAA0330) mp.NA	601120778F1 NIH MGC 20 Homo seplens clink clara MACE: 20072500 mil	Homo sapiens cytochrome P450, subfamily IIF polymentide 4 (Cytopes)	8/88/08.s1 Soares bereithyraid hims NAUDA Homeston	8/88/08.51 Soares barelthyroid times NbHPA Home call	Homo sapiens tumor endothelial marker 7 magnings (TELIA).	Homo saplens tumor endothelial marker 7 precinco (TEMA)DNA	Homo saplens Taxt (human T-cell latikem)s with the District of Taxtons S.	Homo saplens galactocerebrosidase (GALC) gene, exon 15	
	Top Hit Database Source	F	5	Į.	12	LZ	15	Z		Į,	Z	EST HIMAN	FST HIMAN	EST HUMAN	EST HUMAN	EST HUMAN	ST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	Т	HOMAN		EST HUMAN			EST HUMAN		EST HUMAN	HUMAN					
	Top Hit Acession No.	E-85 AF248540 1	202	5174775 NT	5174775 NT	Π	468	Γ	2.0E-85 4505880 NT	4826977 NT	L 163284.2		T				-85 BE618392.1 E					Γ	Γ	-85 AI198420.1 E	11417862 NT	11417862 NT		224	Γ	Γ	88	LN 9889968	11421737 NT	-86 L38557.1 NT	
	Most Similar (Top) Hit BLAST E Value	2.0E-85	2.0E-85	2.0E-85	2.0E-85	2.0E-85	20E-85	2.0E-85	2.0E-85	2.0E-85	20E-85/	2.0E-86	2.0E-85/	2.0E-85		1.0E-85 B	1.0E-85	1.0E-85 B	1.0E-85 A	1.0E-85 A	1.0E-85 B	1.0E-85 B	1.0E-85 Y	1.0E-85 A	1.0E-85	1.0E-85	9.0E-86 BE274217.1	8.0E-86	7.0E-86 A	7.0E-86 AA860801.1	7.0E-86	7.0E-86	7.0E-86	7.0E-86 L3	•
_	Expression Signal	2.34	0.97	8.28	8.28	1.53	5.28	1.18	4.51	1.22	0.97	3.18	1.08	1.32	2.88	8.42	8.42	4.38	2.77	2.77	1.73	1.73	1.28	2.41	4.4	4.74	11.19	1.57	2.34	2.34	1.02	1.02	6.65	3.06	
	ORF SEQ ID NO:	26438	26803	26824	28825	27692		28435	29666	29896	30207	35009	35385	36029		27850	27851	35528	36768	36767	36847	36848	36927	37696	31053	31053	1	37618	26345	26346	31848	31849	30553	34479	
	SEQ ID NO:	13777	14130		14146	•	1	-)	17039	17262	-1 ⁵⁸	21843	22202	22810	15010	15113	15113	22333	23524	23524	23599	23599	23670	24364	24722	24722	4156	24283	13683	13883	18881	18881	17958	21335	
	Probe SEQ ID NO:	1017	1383	1399	1399	2226	2826	3022	8 8 8 8	4527	4854	9173	8549	10162	2285	2382	2362	8	10842	10842	1919	10919	10897	1/13	2020	277	1409	1698	916	916	9403	6103	88	8	

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Тор Нії Descriptar	Homo sepiens RAN binding protein 7 (RANBP7), mRNA	Homo sapiens DiGeorge syndrome critical region gene 6 (DGCR6), mRNA	Homo sapiens similar to transcription factor CA150 (H. sapiens) (LOC63170), mRNA	Homo saplens similar to transcription factor CA150 (H. saplens) (LOC63170), mRNA	Homo saplens oxoglutarate dehydrogenase (lipoamide) (OGDH) mRNA	Homo sapiens enteropeptidase gene, exons 20 and 21	Homo saplens 24 kDa intrinsic membrane protein (PMP24), mRNA	601072594F1 NIH_MGC_12 Homo saplens cDNA clone IMAGE:3458830 5'	601178865F1 NIH_MGC_17 Hamo sapiens cDNA clone IMAGE:3531953 5'	601072594F1 NIH_MGC_12 Homo sapiens cDNA clone IMAGE:3458830 5'	xz92h12.x1 NOL_CGAP_Lu24 Homo saplens cDNA done IMAGE:2871719 3'	AV722329 HTB Hamo sapiens cDNA clone HTBBSD04 5'	601508696F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3911303 5'	601509696F1 NIH_MGC_71 Homo saplens cDNA clone IMAGE:3911303 5'	tu 18b02.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2251371 3'	Homo sapiens myosin X (MYO10), mRNA	EST177232 Jurkat T-cells VI Homo sapiens cDNA 5' end	Homo sapiens chromosome 21 segment HS21C003	yz19808.r1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:283478 5'	Homo sapiens neurexin III (NRXN3) mRNA	Homo sapiens neurexin III (NRXN3) mRNA	Human endogenous retrovinus, complete genome	Homo sapiens mRNA for KIAA1277 protein, partial cds	EST378215 MAGE resequences, MAGI Homo sapiens cDNA	Homo sapiens lysophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	Homo sapiens lysophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	hd87g08.x1 NCI_CGAP_GC6 Homo saplens cDNA done IMAGE:2916542 3'	Homo sapiens cAMP-specific phosphodiesterase &A (PDE&A) mRNA, partial cds	H.saplens mRNA encoding phospholipase c	H.saplens mRNA encoding phospholipase c	Homo sapiens similar to ectonucleotide pyrophosphatase/phosphodiesterase 3 (H. sapiens) (LOC63214),	mRNA	Human Chediak-Higashi syndrome protein short Isoform (LYST) mRNA, complete cds	Hamo saplens chromosome 21 segment HS210027	
Top Hit Database Source	LN FN	ZI.	FN	LN	TN	LN			HUMAN	EST_HUMAN							EST_HUMAN		EST_HUMAN	Ę	Į,						T_HUMAN	Ę	ᅜ	F			NT	NT	
Top Hit Acession No.	5453997 NT	11526307 NT	11417012 NT	11417012 NT	4505492 NT	/19139.1	F005833 NT				1			3.0E-86 BE886479.1		11037056 NT	2.0E-86 AA306264.1		N58977.1	4758827 NT	4768827 NT	9635487 NT		2.0E-86 AW966142.1			1					11419429 NT		2.0E-86 AL163227.2	
Most Similar (Top) Hit BLAST E Value	7.0E-86	7.0E-86	7.0E-86	7.0E-86	6.0E-86	6.0E-86 Y19139.1	6.0E-86	4.0E-86	4.0E-86	4.0E-86	/ 3.0E-86 /	6	3.0E-86	3.0E-86	3.0E-86	3.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2,0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2,0E-86 Z16411.1		2.0E-86	2.0E-86 U84744.1	2.0E-86 A	
Expression Signal	1.39	2.27	1.72	1.72	2.88	2.64	1.07	4.98	12.1	2.18	6.97	1.05	3.37	3.37	5.14	1.6	2.02	2.59	3.21	1.93	1.93	5.09	1.55	1.3	3.54	3.54	2.84	3.26	1.62	1.62		0.69	0.69	0.47	
ORF SEQ ID NO:		35508	36809	36810	26686	30440	30442	25660	31684	25660	31208	33985	35980	35981	36408	37628	25706		26584	26910	26911	27651	27732	28817	29118	.29119		30106	31501	31502		32476	33733		
Exan SEQ ID NO:	22252	22310	23562	23562	14020	17823	17825	13018	18726	13018	18307	20854	22768	22768	23180	24303		13190	13922	14225	14225	14917	14992	16168			16765	17469	18573	18573		25098	20603	21107	
Probe SEQ ID NO:	8696	9858	10882	10882	1271	5105	5107	206	5944	11205	5509	8160	10120	10120	11413	11708	280	405	1168	1478	1478	2188	2266	3410	3729	3729	4018	4737	5782	5782		6974	7908	8414	

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				. 1					l			(98					Γ		IF		F	r		£		1	7	1	1	P	þ	5
	Top Hit Descriptor	Homo sapiens butyrobetaine (gamma), 2-oxoglutarata dioxygenase (gamma-butyrobetaine hydroxylase) (BBOX), mRNA	Homo saplens butyrobetaine (gamma), 2-oxoglutarate dloxygenase (gamma-butyrobetaine hydroxylase)	Homo sanjana ahambalinid	Homo scales of the stranging scrambiase 1 (PLSCR1), mRNA	Home saniens heric helix face Lilling 6.	Homo saplens basic-helix-loop helix-PAS protein (NPAS3), mRNA	Homo sariens menta for Vivia (14 Personal Anno Sarien (NPAS3), mRNA	Homo sepiens mikiya for KiAA1411 protein, partial cds	Homo sapiens through a formation and kinese, 90kD, polypeptide 6 (RPS6KA6) mRNA	Homo sablens dans for AEA complete out	Homo sapitate NADH dehydrogenese (ublquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase)	Homo sapions filtulin 5 (EBI NE) BM.A	Himan damma distant franchistan	Homo septems chambrane 34	Home september the control segment HSZ1C009	Home serious hand-deal	Home conjugate L. C. Control MRNA	Homo saplans chamber of the Colon (LOCo1318), mRNA	Homo conjunctor and the car segment HS21C100	Homo capicals eyilabulgalin 1 (OYNO1), mKNA	Homo septence chromosom 21 segment HSZ1C084	9b77c09x1 Soares, Edal heart, NbHH19W Homo saplens oDNA clone IMAGE:1706128 3' similar to	DIVISION OF THE PUZZES REPAIN, TYPE CYTOSKELETAL 10	Home sapiens a disintegrin and metalloproteinase domain 22 (ADAM22), mRNA	month septems a distintegrin and metalloproteinase domain 22 (ADAM22), mRNA	O.Cuniculus mRNA for elongation factor 1 alpha	// NSDIVEXTINCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:3322779 3'	Institution CCAP Co16 Homo sapiens cDNA clone IMAGE:3322779 3'	MRC-N 10039-020500-004-a11 NT0039 Homo sapiens cDNA	IL3-H10619-060700-198-D10 HT0619 Homo sapiens cDNA	LS-H10702-160600-103-d06 HT0702 Homo sapiens cDNA
	Top Hit Database Source	FZ.	Ę	Į	Į.	ΙN	Ļ	LZ	Į.	5	¥	<u> </u>	Į	LN L	Į	Ę	5		; <u> </u>	; ;	Ę	Ę	TOT LIMAN	-1.			- N	EST HUMAN	EST HOMAN	NAMON TO	ESI HUMAN	EST HOMAN
	Top Hit Acession No	11437135 NT	11437135INT	10863876 NT	11422084 NT	11545848 NT	11545846 NT	86 AB037832.1	4759051 NT	11418189INT	86 AB011399.1	9855	5453649 NT		1.0E-86 AL163209.2		19181	7708184 NT	L163300.2	1.0E-86 4507334 NT	1.0E-86 AL 163284 2			E77794	TN 127724	Vegade 4		T	T,	T	T	
	Most Similar (Top) Hit BLAST E Value	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-86	2.0E-88	2.0E-86	1.0E-86	1.0E-86	1.0E-86 L20492.1	1.0E-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86 A	1.0E-86	1.0E-88 A	1.0E-86 A	9.0E-87.A	9 0F-87	9.0E-87		7 70 = 87 B	7 OF-87 BS		7 0E-87 BE		
	Expression Signal	2.31	2.31	0.65	1.95	2.91	2.91	1.15	2.54	3.07	4.26	2.28	1.5	2.7	1.32	1.32	0.88	0.88	5.2	1.23	2	1.37	1.81	1.7	1.7	15.93	1 79	1 70	0.57	3.4	4 45	
	ORF SEQ ID NO:	34307	34308	34840	35067	36210	36211	36262	36742	31027		27027	28569	28645	28699	28700	29318	29319	29599	29941	31155	31155		33105	33106	25893	27755	27756	32064	33913	33544	
	SEQ ID NO:	21164	21164		21897	22892	22992	23048	23503	24827	24926	14338	15923	15992	16051	16051	16676	16676	16974	17313	18264	18264	18078	2002	2002	13252	15019	15019	19079	20783	20425	
	Probe SEQ ID NO:	8472	8472	8801	9218	10345	10345	10400	10820	12458	12621	1592	3160	3229	3280	3290	3926	3926	4233	4578	5465	11606	6272	7348	7348	467	2294	2294	6307	6808	ı	

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Single Exon Probes Expressed in brain	ORF SEQ Expression (Top) Hit Acesskon Database ID NO: Signal BLAST E No. Source	AL043314.2	3.85 7.0E-87 AL043314.2 EST_HUMAN	7.0E-87 AI081565.1 EST_HUMAN	6.65 7.0E-87 K03002.1	7.0E-87 K03002.1 NT	6.0E-87 7657213 NT	0.69 6.0E-87 7657213 NT	32085 2.02 6.0E-87 AB029004.1 NT Home Septembring to recommend the comment of the		5.0E-87 AA382811.1 EST_HUMAN	1.56 5.0E-87 AA382811.1 EST_HUMAN	1.51 4.0E-87 AL163210.2 INT	13.58 4.0E-87 AB037835.1 NT	4.0E-87 AB007925.1 NT	1.03 4.0E-87 7706299 NT	1.03 4.0E-87 7706299 NT	TO TO TO TO	TOGOSOLWO	2.77 4.0E-87 COUSZ1 SW155FRC1	T HUMAN	0.72 4.0E-87 L48524.1	7 M60676.1 NT	1.5 4.0E-87 11417862 NT	1.5 4.0E-87 11417862 NT	2.25 4.0E-87 11417812 NT	2.0E-87 4885420 NT	0.83 2.0E-87 AU116935.1 EST_HUMAN	2.0E-87 BF376311.1 EST_HUMAN	1.47 2.0E-87 BE175478.1 EST_HUMAN	2.0E-87 BE734190.1 EST HUMAN	2.0E-87 BE734190.1 EST_HUMAN	9.81 2.0E-87 BE567193.1 JEST_HUMAN 001341385F1 NIT MCC_35 TOLIN SQUARE CONTROLL
																					31695		37065	30721				29154	30209				
	Exon SEQ ID NO:	22619	22619			I _		i i	\	23318	1	L	13711	L	L	L	┸	1		18162	18736	1_	L	L	<u> </u>	L		16516	<u>L</u> .	L	5 18372	ľ	1 1
	Probe SEQ ID NO:	9971	9971	10366	10806	10806	3517	5128	6327	10625	1135	12207	ğ	11/40	2024	2424	242	34.7	3457	5360	5054	7670	11118	12396	12396	12541	2779	3764	4857	4907	557	5275	6234

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		wodant d Seemed in the	1972 IOV.: 1 Soares retai (iver spieen 1NFLS Homo saplens cDNA clone IMAGE:243396 5	AV634143 GLC Hamo sapiens cDNA clone GLCDSG04 3'	BUTT/BU3ZF1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3531511 6	Homo sapiens hect domain and RLD 2 (HERC2), mRNA	WZ1607.r1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:243398 61	yv21607.r1 Scares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:243386 5	Fruman cydophillin gene for cyclophilin (EC 5.2.1.8)	0012/8312F1 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3810539 6'	Para Septets putative glycolipid transfer protein (LOC51054), mRNA	FMZ-C 10285-141099-001-g04 CT0285 Homo saplens cDNA	Historio Locabe 141089-001-904 C10265 Homo sapiens cDNA	Home coning the recommendation of the recomm	Home septiens neurean III (NRXN3) mRNA	Home septens intersectin long isoform (ITSN) mRNA, complete cds	ndrino sepiens intersectin long isoform (ITSN) mRNA, complete cds	Home carloss and the same appears con cone IMAGE:1620199 3'	Homo saniens asserted factor receptor-bound protein 10 (GRB10) gene, exon 8	Homo serviens confliction:	Homo serviers conflictional	Homo saplens 10 modifications of the company of the	Homo sablens profess Kingao C Letter (April 2017) Homo sablens profess (Index Professor)	Homo caniene trachol out.	Homo septens mRNA for although a city.	domo saplena mRNA for altaba 3 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	ROBBN0278-05000000 Box BN032-011.	DOS BNOSTS SESTING 12-EUZ BNOZ/6 Homo saplens cDNA	KCo-BNUZ/6-060700-012-E02 BN0278 Homo saplens cDNA	ruman L-plastin mRNA, 5'end	name saplens nect domain and RLD 2 (HERC2), mRNA	Tromo sapiens RGH1 gene, retrovirus-like element	Home saviers Editive repeats and discoldin I-like domains 3 (EDIL3), mRNA	Trumo septens Edir-like repeats and discoldin I-like domains 3 (EDIL3), mRNA	ndno sapiens suitotransferase-related protein (SULTX3), mRNA	nono sapiens sulfotransferase-related protein (SULTX3), mRNA
igle Exon Pro	Top Hit Database Source		FOT HIMAN	NUMBER TO LOCAL	EST HOMAN	NAMOR I	IN LOL	EST HUMAN	ESI_HOMAN	TOT TOTAL	ESI TOMAN	FOT LIMAN	EST HIMAN	L L		F		FST HIMAN	LN LN	NT.				ľ	Į.			T HUMAN	T	NAMOL							
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	Expression Signal		0.69	0.81	1.58	0.94	39.61	35.45	17.42	5.72	2.09	1.1	1-1	6.23	2.43	0.69	0.69	1.04	1.91	1.91	0.62	0.62	1.18	1.23	12.83	0.97	0.97	2.85	2.85	0.67	1.65	1.92	1.68	1.68	3.54	3.94	
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	Top Hit Descriptor	Hown sapiens double stranded RNA activated protein kinase (PKR) gene, exon 12	Line contact m DNA for KIAA 1399 profess cases	Home suprementation in Annual to programs	Hano sapiens many tor NIAA 1309 protein, paren 3000	Homo saplens DKF-2P686F1322 protein (Dixt 2F 900) 1022, marks	Hamo sapiens chromosome ZI segment HSZ I Cours	H.sapiens ECE-1 gene (excn 9)	H.sapiens ECE-1 gene (excn 9)	Homo saplens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, CACLES, CAC	Homo cantans X-linked anhidrolitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	Homo sapiens KIAA0063 gene product (KIAA0063), mixwa	Kg719F Human fetal heart, Lambda ZAP Express Homo sapiens curva cione no 18.5 similar. Service FINGER PROTEIN HZF1	Homo sapiens Intersectin short isoform (ITSN) mRNA, complete cds	Wd68h08,X1 NCI_CGAP_Lu24 Home saplens cUNA done IMAGE 23307.99 5 3111161 (3 5 cm. 1	repetitive element;contains element MER22 MER22 repetitive element;	Horno Squeris lines econ driver the Homo Sapiens cDNA clone IMAGE:47129 5	Wildon's Chromosome 21 segment HS21C084	ANSTALOFRET NIH MIGG 83 Homo sapiens cDNA clone IMAGE:4295775 5	DX41-TN0028-050900-004-f10 TN0028 Homo sapiens cDNA	PMA-TN0028-050900-004-f10 TN0028 Homo saplens cDNA	Home saplens transforming growth factor, beta-induced, 68kD (TGFBI), mRNA	Homo sapiens cell division cycle 10 (homologous to CDC10 of S. cerevisiae) (CDC10) mRNA	Hamo sapiens KIAA0152 gene product (KIAA0152), mRNA	Homo sapiens KIAA0152 gene product (KIAA0152), mRNA	Homo sapians hypothetical protain FLI21634 (FLI21634), mRNA	Homo sabiens zinc finger protein 259 (ZNF259) mRNA	za48112.51 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:285823 3'	Homo sariens a disintegrin and metalloproteinase domain 23 (ADAM23) mRNA	Homo sapiens a disintegrin and metalloproteinase domain 23 (ADAM23) mRNA			
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	Top Hit Acession No.	T			4B037820.1	7661701 NT	4L163209.2		X91929.1		AB026898.1	AF003528.1	7661887 NT	Nieozog 4	AF114488.1		AI693217.1	AF114488.1	H10932.1	AL163284.2	BF680206.1	BF091229.1	BF097229.1	AEOSEO NT	450/2094 IN				10001	Noosol. I			11429300 NI	
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	Expression Signal		10.24	2.78	2.76	1.57	1 35	273	27.0		1.05	3.82	100				2.28	0.83	3.19	1.8														2.85
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1 **40667** Homo saplens molybdenum cofactor blosynthesis profein A and molybdenum cofactor blosynthesis protein Homo sapiens growth differentiation factor 6 (cartilago-derived morphogenetle protein-1) (GDF6), mRNA wq70a12.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2476606 3' aa64a11.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:824732 3' similar to WP:B0272.2 zp87c02.r1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone IMAGE:627170 5' similar to SW:POL1_HUMAN P10266 RETROVIRUS-RELATED POL POLYPROTEIN; DKFZp434N0323_J1 434 (synonym: htes3) Homo saplens dDNA clone DKFZp434N0323 5° os91g03.s1 NCI_GGAP_GG3 Homo saplens dDNA clone IMAGE:1612766 3° similar to gb:M16342 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEINS C1/C2 (HUMAN); Homo sapiens Calsenilin, presenilin-binding protein, EF hand transcription factor (CSEN), mRNA Homo sapiens SNARE protein kinase SNAK mRNA, complete cds UI-H-BIT-aea-d-04-0-UI.s1 NCI_CGAP_Sub3 Homo saplens cDNA clone IMAGE:2718750 3' UI-H-BIT-aea-d-04-0-UI.s1 NCI_CGAP_Sub3 Homo saplens cDNA clone IMAGE:2718750 3' Homo sapiens v-ral simian leukamia viral oncogene homolog A (ras related) (RALA), mRNA Homo saplens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA Homo saplens mRNA for RALDH2-T, complete ods Homo saplens acyt-Coenzyme A dehydrogenese family, member 8 (ACAD8), mRNA Homo saplens transcobalamin II; macrocytic anemia (TCN2), mRNA Homo sapiens polycythemia rubra vera 1, cell surface receptor (PRV1), mRNA Homo sapiens SNARE protein kinase SNAK mRNA, complete cds Homo sapiens dynein, axonemal, light polypeptide 4 (DNAL4), mRNA Homo sapiens SNARE protein kinase SNAK mRNA, complete cds Homo sapiens retinoblastoma-binding protein 2 (RBBP2), mRNA Top Hit Descriptor Homo sapiens putative anion transporter 1 mRNA, complete cds Homo sapiens activator of S phase kinase (ASK), mRNA Homo sapiens activator of S phase kinase (ASK), mRNA Homo saplens mRNA for RALDH2-T, complete cds Homo sapiens chromosome 21 segment HS21C046 Homo sapiens KIAA0417 mRNA, complete cds Homo saplens KIAA0417 mRNA, complete cds Homo saplens Interfeukin 13 (IL13), mRNA Single Exon Probes Expressed in Brain aa54a11.s1 NCI_CGAP CE00851; mRNA, complete cds Top Hit Database HUMAN Source EST_HUMAN HUMAN EST_HUMAN HUMAN 11417370 NT 11419210 NT EST_ EST 11421726 NT 눋 11419210 NT 11436400 NT 998688 Fop Hit Acession 11420697 7305198 11439065 11417974 2.0E-88 AF246219.1 2.0E-88 5031666 AF279265.1 AF034374.1 2.0E-88 730 2.0E-88 AF246219.1 .0E-88 AW139565.1 AB015228.1 AF246219.1 1.0E-88 AA488981.1 AL043314.2 1.0E-88 AW139565. AB007877.1 1.0E-88 AA190368.1 1.0E-88 AA991479.1 1.0E-88 AL163246.2 1.0E-88 AI969034.1 3.0E-88 3.0E-88 3.0E-88 3.0E-88 3.0E-88 3.0E-88 3.0E-88 3.0E-88 Most Simila (Top) Hit. BLAST E 3.0E-88 3.0E-88 3.0E:88 0E-88 2.0E-88 9.6 6.15 9.58 1.28 3.13 2.13 1.52 Expression 5.83 23.81 4.42 0.6 2.99 Signal 2.97 ORF SEQ ID NO: 31812 32076 32077 32465 33642 33206 33921 33521 35680 27185 28867 29760 31539 35681 26432 27056 35705 32317 32839 35037 36420 35311 18296 18848 25088 SEQ ID 25088 20516 19447 20117 22479 13773 14367 16214 18610 19313 19313 19774 23189 ÿ 21872 24760 22131 8 Probe SEQ ID 6319 6319 5618 6965 7821 9334 9828 9828 1620 1744 3458 9867 1013 5821 5821 6548 8548 7022 ÿ 9141 4391 7084 11422 12356

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| 801142409F1 NIH_MGC_14 Homo sapiens cDNA clane IMAGE:3506186 5" | Homo sepiens similar to sema domain, immunoglobulin domain (Ig), short basic domain, secreted, (semaphorin) 3A (H. sepiens) (LOC63232), mRNA | Homo sapiens hormonally upregulated neu fumor-associated kinase (HUNK), mRNA | Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA | Homo saplens complement component 8, beta polypeptide (C8B) mRNA | DKFZp434E246_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434E246 5' | H.saplens CLN3 gene, complete CDS | H.saplens CLN3 gene, complete CDS | Homo sepiens plastin 3 (T isoform) (PLS3), mRNA | Homo sapiens plastin 3 (T isoform) (PLS3), mRNA | Homo sapiens actin related protein 2/3 complex, subunit 1A (41 kD) (ARPC1A), mRNA | Homo sapiens KIAA0433 protein (KIAA0433), mRNA | Homo sapiens KIAA0433 protein (KIAA0433), mRNA | Homo sapiens mRNA for KIAA0561 protein, partial cds | H.sapiens Wee1 hu gene | H.sapiens Weet hu gene | Homo sapiens mRNA for KIAA0823 protein, partial cds
 | Homo sapiens mRNA for KIAA0823 protein, partial cds | Human aldose reductase (AR) gene, segment 2 | Human aconitate hydratase (ACO2) gene, exon 2 | Homo sapiens inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA
 | Homo sapiens serine/threonine-protein kinase PRP4 homolog (PRP4) mRNA
 | Homo sapiens ubiquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA | Homo sapiens ubiquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA | Homo sapiens HSPC159 protein (HSPC159), mRNA
 | Homo sapiens mRNA for KIAA0406 protein, partial cds | Homo sapiens mRNA for KIAA0406 protein, partial cds | Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA | Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA | TCBAP2E0383 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sa
 | cDNA clone TCBAP0383 | TCBAP2E0383 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo se
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| 1,33 | 1.2 | 1.72 | 1.72 | 2.86 | 3.35 | 1.34 | 1.34 | 0.57 | 0.57 | 2.06 | 0.57 | 0.57 | 9.0 | 1.11 | 1.11 | 2.33
 | 2.33 | 1.45 | 1.7 | 6.73
 | 1.27
 | 1.06 | 1.06 | 0.88
 | 3.02 | 3.02 | 0.81 | 0.81 |
 | 2.74 | 27.6 | |
| 28184 | 32519 | | 25857 | 30181 | 30228 | 30827 | 30828 | 31997 | 31998 | 33156 | 33598 | 33599 | 35421 | 36296 | 36297 | 36320
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| 15445 | 19495 | 13210 | 13210 | 17559 | 17605 | 18148 | 18148 | 19024 | l i | 20076 | 20474 | 20474 | 22237 | 23075 | 23075 | 23091
 | 23091 | 23867 | 25028 | 13762
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| 2739 | 6833 | 424 | 424 | 4828 | 4878 | 5345 | 5345 | 6250 | 6250 | 7398 | 7779 | 7779 | 9584 | 10429 | 10429 | 10445
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 | 4593 | 4593 | 5100 | 5100 |
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| | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.772 7.0E-89 7557213 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7557213 NT 13210 25857 1.72 7.0E-89 7557213 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17559 30181 2.86 7.0E-89 4557390 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17559 30181 2.86 7.0E-89 4557390 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7687213 NT 13210 25857 1.72 7.0E-89 7687213 NT 17559 30181 2.86 7.0E-89 4557390 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X89832.1 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17559 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X99832.1 NT 18148 30828 1.34 7.0E-89 X99832.1 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17559 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X59832.1 NT 18148 30828 1.34 7.0E-89 X59832.1 NT 19024 31997 0.57 7.0E-89 7549808 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25886 1.72 7.0E-89 7657213 NT 1759 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 A557360 NT 18148 30827 1.34 7.0E-89 X99832.1 NT 18024 31997 0.57 7.0E-89 X549808 NT 19024 31998 0.57 7.0E-89 7549808 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17569 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X59832.1 NT 18148 30828 1.34 7.0E-89 X59832.1 NT 19024 31997 0.57 7.0E-89 7549808 NT 19024 31998 0.57 7.0E-89 7549808 NT 20076 33156 2.06 7.0E-89 7549808 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17569 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X98832.1 NT 18148 30828 1.34 7.0E-89 X98832.1 NT 19024 31997 0.57 7.0E-89 7549808 NT 19024 31998 0.57 7.0E-89 7549808 NT 20076 33156 2.06 7.0E-89 11420754 NT 20474 33598 0.57 7.0E-89 11417118 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 17569 30181 2.86 7.0E-89 4557360 NT 17605 30228 3.35 7.0E-89 AL045748.1 EST_HUMAN 18148 30827 1.34 7.0E-89 X98832.1 NT 18149 30828 1.34 7.0E-89 X98832.1 NT 18024 31997 0.57 7.0E-89 7549808 NT 19024 31998 0.57 7.0E-89 7549808 NT 20076 33156 2.06 7.0E-89 11447118 NT 20474 33599 0.57 7.0E-89 11447118 NT | 15445 28184 1,33 8,0E-89 BE311557,1 EST_HUMAN 19495 32519 1,2 8,0E-89 11421514 NT 13210 25866 1,72 7,0E-89 7657213 NT 1750 30181 2,86 7,0E-89 7657213 NT 17659 30181 2,86 7,0E-89 4557360 NT 17605 30228 3,35 7,0E-89 4557360 NT 18148 30827 1,34 7,0E-89 7569832.1 NT 18024 31997 0,57 7,0E-89 7564980 NT 19024 31998 0,57 7,0E-89 7564980 NT 20076 33156 2,06 7,0E-89 7549808 NT 20474 33599 0,57 7,0E-89 11417118 NT 20474 33599 0,57 7,0E-89 11417118 NT 20474 33599 0,57 7,0E-89 11417118 NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 19495 32519 1.2 8.0E-89 11421514 NT 13210 25886 1.72 7.0E-89 7657213 NT 1759 30181 2.86 7.0E-89 7657213 NT 17659 30181 2.86 7.0E-89 7657213 NT 17605 30228 3.35 7.0E-89 4557360 NT 18148 30827 1.34 7.0E-89 X98932.1 NT 18024 31998 0.57 7.0E-89 7549808 NT 19024 31998 0.57 7.0E-89 7549808 NT 20076 33156 2.06 7.0E-89 7549808 NT 20474 33599 0.57 7.0E-89 11417118 NT 22237 35421 0.6 7.0E-89 NB011133.1 NT 22075 362048.1 1747718 NT NT | 15445 28184 1.33 8.0E-89 BE311557.1 EST_HUMAN 13210 25856 1.72 7.0E-89 7657213 NT 13210 25857 1.72 7.0E-89 7657213 NT 1759 30181 2.86 7.0E-89 7657213 NT 17659 30181 2.86 7.0E-89 7657213 NT 17605
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Page 398 of 536 Table 4 Single Exon Probes Expressed in Brain

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Single Exon Probes Expressed in brain	Top Hit Descriptor	QV3-NT0022-080600-219-g03 NT0022 Homo saplens cDNA	we91c03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2348452.37	EST388290 MAGE resequences, MAGN Homo sapiens cDNA	qh17b06.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1844915 3'	Human mRNA for KIAA0299 gene, partial cds	w88e11.r1 Soares_placenta_8to9weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:259148 6' similar to SW:PI4K_HUMAN P42356 PHOSPHATIDYLINOSITOL 4-KINASE ALPHA;	Homo sapiens PXR2b protein (PXR2b), mRNA	Homo sapiens PXR2b protein (PXR2b), mRNA	Homo saplens PXR2b protein (PXR2b), mRNA	Homo sapiens PXR2b protein (PXR2b), mRNA	Homo saplens mRNA for KIAA1342 protein, partial ods	qg98c08.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1843022.3' similar to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);contains Alu repetitive element;	ah70e03.s1 Soares_testis_NHT Homo sapiens cDNA clone 1320988 3'	ah70e03.s1 Soares_testis_NHT Homo sapiens cDNA clone 1320988 3'	Homo sapiens topoisomerase-related function protein (TRF4-2) mRNA, partial cds	H.saplens HCK gene for tyrosine kinase (PTK), exons 10-11	H.saplens HCK gene for tyrosine kinase (PTK), exons 10-11	Homo sapiens chromosome 21 segment HS21C003	Homo sapiens GGT gene, exon 5	601065986F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3452423 5	Homo sapiens gene for LECT2, complete cds	Human N-ethylmaletmide-sensitive factor mRNA, partial cds	Homo sapiens chromosome 21 segment HS21C085	Human GT24 (GT24) mRNA, partial ods	Homo sapiens solute carrier family 24 (sodium/potassium/caldium exchanger), member 2 (SLC24A2), mRNA	Homo sapiens partial mRNA for PEX5 related protein	Homo saplens mRNA for KIAA1333 protein, partial cds		Homo sapiens CaBP5 (CABP5) gene, exon 5
JIE EXOII PIO	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	FN	EST_HUMAN	Į.	F	FZ	ZI.	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝT	TN	NT	NT	NT	EST_HUMAN	Ę	TN	NT	NT	۲	LN LN	LN.	TN	NT
Silio Silio	Top Hit Acession No.	4.0E-89 BE762749.1		l		•		7706670	T706670 NT	TN06870 NT	TN06670 NT	89 AB037763.1	89 AI222095.1	89 AA759149.1		89 AF089897.1			89 AL163203.2	89 AJ007378.1	89 BE541744.1	1.1	89 U03985.1	.2	89 U81004.1	11428801	89 AJ245503.1	89 AB037754.1	-89 AF170814.1	
	Most Similar (Top) Hit BLAST E Value	4.0E-89	4.0E-89	3.0E-89	3.0E-89	3.0E-89	3.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89	2.0E-89
	Expression Signal	1.85	1.56	1.51	1.28	0.48	2.34	0.87	0.87	1.55	1.55	3.17	1.53	1.01	1.01	1.26	5.16	5.16	0.75	1.1	99.0	2.9	1.61	0.63	5.33	3.07	1.02	9.0	0.68	0.68
	ORF SEQ ID NO:	33251	37034		32790	36374	36630	25832	25833	25832		25933	28293	28946	28947	29494			29679	29825		30886	31418	31861	33341	33657	34146	35052	36657	35558
	SEQ ID NO:	20159	23758	<u> </u>	19731	23148	23393	13184	13184	13184	13184	13301	15650	16296	•	1	16875	16875	17054	17199	18065	18193	18496	1	20237	20530		ı	22361	l i
	Probe SEQ ID NO:	7487	11088	2879	7040	10502	10702	123	123	88	386	517	2883	3540	3540	4125	4133	4133	4315	4463	5269	5393	5702	6116	7667	7835	8316	9152	9710	9710

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	Top Hit Descriptor	Homo saplens integrin, alpha 3 (antigen CD49C, alpha 3 subunit of VLA-3 receptor) (ITGA3), mRNA	Homo sapiens hect domain and RLD 2 (HERC2), mrNA	Homo saplens cell adhesion molecule with homology to L1CAM (close homologue of L1) (CHL1), mRNA	Human MAGE-7 antigen (MAGE7) pseudogene, complete cds	hre1d09.X1 NCI_CGAP_Kid11 Hamo sapiens cDNA clane IMAGE:3134697 3 similar to 11.00-11.00	SOLUTE CARRIER FAMILY 22-LIKE 2 PROTEIN ; Instance ANCI CGAP Kid11 Homo sepiens cDNA clone IMAGE:3134897 3' similar to TR:O54778 O54778 Instance ANCI CGAP Kid11 Homo sepiens cDNA clone IMAGE:3134897 3' similar to TR:O54778 O54778	SOLUTE CARRIER FAMILY 22 - LIKE 2 PROTEIN ;	Homo sepiers difference 21 cognition HS21C046	Homo sapiens circumsome 21 segment HS210046	Homo sapiens circumscane 21 segment HS21C046	Homo saptems circulations and a construction of the construction o	7636f08.X1 NCI_CGAP_Luz4 nonito septents 0011.	7636708 X1 NCL CGAP Lu24 Homo sapiens con concentrations	RC1-HT0598-120400-022-b08 HT0598 Homo sapiens cDNA		qg98c08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1843022 3' similar to gb:JU4131 GBAMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);contains Alu repetitive element;	(AAC) - 4m ed and (am) - 4m ed ed ed ed ed ed ed ed ed ed ed ed ed	qg96c08.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1843/022 3 similar to go.ov-roll gaMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);contains Alu repetitive element; agaMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);contains Alu repetitive element; and a saplens cDNA clone IMAGE:461442 3	282g10.s1 Soares letal live spread ANEIS S1 Homo seplens cDNA clone IMAGE:461442 3'	282g10.s1 Soares fetal liver spired 114 20 and partial cds, alternative (CACNA1F) gene exons 7-49, and partial cds, alternative (CACNA1F) gene exons 7-49, and partial cds, alternative (CACNA1F) general exons 7-49, and partial cds, alternative (CACNA1F) general exons 7-49, and partial cds, alternative (CACNA1F) general expensive (CACNA1F	Homo sapiens calcium channer aprila in sacering (construction)	spliced cares tests NHT Homo sapiens cDNA clone 1375503 3'	ANAESS837R1 NIH MGC 66 Home septens cDNA clone IMAGE:3855824 3	AN 6555R37R1 NIH MGC 66 Homo sapiens cDNA clone IMAGE:3855824 3	vr88e04.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:212190 3' similar to	SP.C1TC_HUMAN P11586 C-1-TETRAHYDROFOLATE SYN I HASE, CT OF LASMING.	SPICITIC HUMAN P11586 C-1-TETRAHYDROFOLATE SYNTHASE, CYTOPLASMIC;	
Siligie Exoli r 10083	Top Hit Database Source	·					EST_HUMAN	EST_HUMAN	N	IN	Z.		EST HUMAN	EST HUMAN	EST HUMAN		EST HUMAN		EST_HUMAN	EST HUMAN	EST_HUMAN		LN L	ES! HUMAN	TOT TOTAL	NEW TOP I SE	EST_HUMAN	EST_HUMAN	
Billo	Top Hit Acession No.	11434411 NT	5729867 NT	11433673 NT	140602 4	110092.1	BF196052.1	BF196052.1	AL163246.2	AL163246.2	AL163246.2	AL163246.2	BE670561.1	DER70561 1	DEG/0301.1	BE17 / 020. 1	0.12220055.1	AIZZZUBO. I	A)222095.1	AA705222.1	AA705222.1		AF223391.1	0 AA782977.1	7.0E-90 BE962525.2	BE962525.2	7.0E-90 H68849.1	7.0E-90 H68849.1	
	Most Similar (Top) Hit BLAST E Value	2 OF-89	201.0	00			1.0E-89	1.0E-89	9.0E-30	9.0E-90		8.0E-90	8.0E-90	20.0	8.05-90	8.05-90	11 0	8.05-90	8.0E-90	8.0E-90	8.0E-90		7.0E-90						
	Expression Signal	684	3 6		5.03	2.11	5.97	5.97	1.67	1.57	2.23	2.9	3.78	2	3.78	0.55	•	1.52	1.52				4.12	2.08		1.82	1.9	9	
	ORF SEQ ID NO:	-	37339	\perp		37636	37483	37484	L	1						34293		36531	36532	١				100	34701		35901		32902
	SEQ ID		24036	C1282	24161	24312	24169	24160	Canc		L	1	ᆚ			21150	<u> </u>	23293	20000	1	1	8007	13587		L	ı	22684	ı	22684
	Probe SEQ ID		1348	11448	11562	11718	11570	44670	0,100	3 6	2 15	100	1047	1307	1307	8458		10599		SEOL.	10963	10903	816	8323	88	8865	10036	3	10036

Page 400 of 536 Table 4 Single Exon Probes Expressed in Brain

ı		Т	T	Т	Т	Т	Т	Т	T	Т	т —		1	1	Т	т-	Ti	7-1			<u>.</u>	11 4	W.,	1	Tř	11 -		16	· ·	75 17	J
		L		l		l	ı	İ			3	3-			İ	L	"	ľ	T	ľ	ĺ	ľ-	۲	1	*	٦-		T.	1	1	567
Ongo ryprosed in Dall	Top Hit Descriptor	602071208F1 NCI_CGAP_Bm64 Homo sapiens oDNA clone IMAGE:4214257 5'	Homo sapiens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo sapiens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo seplens HsGCN1 mRNA, partial cds	Homo sapiens HsGCN1 mRNA, pertial ods	Homo saplens inositol 1,4,5-triphosphate receptor, type 3 (ITPR3) mRNA	Homo sapiens inositol 1,4,5-triphosphate receptor, type 3 (ITPR3) mRNA	Homo sapiens TCL6 gene, exon 1-10b	Human gamma-aminobutyric acid transaminase mRNA, partial cds	999608.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1843022 3' similar to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);contains Alu repetitive element;	999608.xt Soares, NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:1843022 3' similar to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN):contains Alu repetitive element:	Homo saplens Intersectin long Isoform (ITSN) mRNA, complete cds	Homo saplens pregnancy-zone protein (PZP) mRNA	DKFZp762P1616_11 762 (synonym: hmel2) Homo sapiens cDNA clone DKFZp762P1616 6'	H.sapiens mRNA encoding phospholipase c	Homo sapiens ELKS mRNA, complete cds	H. sapiens mRNA encoding phospholipase c	Homo saplens Carbonic anhydrase-related protein 10 (LOC56934), mRNA	Homo saplens Carbonic anhydrase-related protein 10 (LOC56934), mRNA	Homo saplens angiopoietin 4 (ANG4) mRNA, partial cds	Homo saplens anglopoletin 4 (ANG4) mRNA, partial cds	Homo saplens adenylate cyclase 9 (ADCY9) mRNA	Homo saplens ryanodine receptor 3 (RYR3) mRNA	Homo saplens hypothetical protein FLJ13222 (FLJ13222), mRNA	Homo saplens cadherin 18 (CDH18) mRNA	Homo saplens similar to ectoruoleotide pyrophosphatase/phosphodiesterase 3 (H. saplens) (LOC63214),	Homo contana radri im hindina fromenador mDNA marini ada	Home emiser KIA 60499 amelia (KIA 6049) amelia (KIA 6049) amelia (KIA 6049) amelia (KIA 6049) amelia (KIA 6049)	Homo septems Nitherous protein (Nitheotas), IIINNA Homo septems (National and and Alas), IIINNA	Charles (Corporate) install (Corporate) install corporate single outside outsi
יייי בייייי פוני	Top Hit Database Source	EST_HUMAN	۲	N L	Ę	LN TN	N	۲	NT	NT	EST_HUMAN	EST HUMAN	NT	NT	EST HUMAN	۲	N	NT	LN	NT	LN	NT	NT	NT	NT	NT	Ę	12			
5	Top Hit Acassion No.	BF526089.1	8922398 NT	8922398 NT	90 U77700.1	6.0E-90 U77700.1	4504794 NT	4504794 NT	90 AB035344.1	90 U80226.1	90 AI222085.1	AI222095.1	5.0E-90 AF114487.1	4506354 NT	6.0E-90 AL135549.1		5.	216411.1	9910365 NT	9910365 NT	5.0E-90 AF113708.1	5.0E-90 AF113708.1	4557258 NT	4506758 NT	11345483 NT	4826670 NT	14 00707777	275	6 OF OO 44447449 NIT	TN 4177444	
	Most Similar (Top) Hit BLAST E Value	7.0E-90	6.0E-		6.0E-		١.	6.0E-90	5.0E-90	5.0E-90	5.0E-90	5.0E-90	6.0E-90	6.0E-90	6.0E-90	5.0E-90	6.0E-90	5.0E-90	5.0E-90	5.0E-90	5.0E-90	5.0E-90	5.0E-90	6.0E-90	5.0E-90	6.0E-90	00 10 2	2.0E-90	00 10 4	90.0	
	Expression Signal	1.17	9.12	9.12	3.27	3.27	2.75	2.75	19.84	3.08	1.47	1.47	2.79	2.05	0.98	2.94	1.31	2.36	0.74	0.74	2.04	2.04	10.6	0.44	5.08	0.58	90	200	200	200	
	ORF SEQ ID NO:	36216					34056			26586	. 27267	27.268	28011	29871	29992	31203	31316	31203	32407	32408	32866	32867	33228	33609	34025	34950	95446	36044	38475	38178	
	Exon SEQ ID NO:	22999		16942		18679		20919		13924	14553	14553	15274	17238			18402				- 1	19802	- 1	20485		21784	COCCC		Л.	22050	1
	Probe SEQ ID NO:	10352	4201	4201	5894	5894	8225	8225	151	1170	1813	1813	2560	4603	4622	5504	5606	5679	6631	6631	7114	7114	7463	7790	8192	9096	0630	10181	10342	10312	

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			Т	Т	Т	T	Т	1		Г	Г	Γ	Γ	Г	Ţ	Г	Γ	Г	T P		1	Γ		U E	3	I],.			SE
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens ATPase, aminophospholipid transporter-like, Class I, type 8A, member 2 (ATP8A2), mRNA	Homo sapiens KIAA0317 gene product (KIAA0317), minva	Homo sapiens KIAA0317 gene product (KIAA0317), mrvva	Homo sapiens KIAA0305 gene product (KIAAU300), mrkva	Homo sepiens gene for AF-6, complete cds	ar/8h05,X1 Barstead adria nin-line appeals with design in the control of the cont	Homo sapiens chromosome 21 univident mRNA	Homo septens circumosome z i convictori in contra de la contra del la contra de la contra de la contra del la contra del la contra de la contra de la contra del la con	Home squere injusting discoulding scenarior bankingse, exon 16	H. saperis gerie directing discount receptor (PSM) gene, complete cds	Homo saprens prostate specific mental and analysis (1972) mRNA	Homo sapiens low general propriational process (2 (2 2 2)	Homo sepiens fow density lipoprotein-realted proteins (12 4 7) missing in the contract of the	Homo sapiens DINA for amyida predurad protein, cariptere cas	Homo saplens mixing for NIPA Lett protein, partein coo	Human promorting to a little And Code Subst Homo samens con Adde Good IMAGE:3083839 3	UI-H-BW T-BM - AND THE A NOT COAP Sub7 Homo segiens oDNA clone IMAGE:3083639 3'	101-H-50W 1-Bity-0-4-CO. 101-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0	601333244F1 Nin MCC 39 Home sapers CDNA clone IMAGE:3453834 5	United services high-mobility group (nonhistone chromosomal) protein 17 (HMG17), mRNA	Home sepiens high-mobility group (nonhistone chromosomal) protein 17 (HMG17), mRNA	Annual Control of Science Information Biogeneeks 2NbHP8to9W Homo septens cDNA clone IMAGE:1713410 3	similar to SW: OLF3_MOUSE P23275 OLFACTORY RECEPTOR OR3.;	Homo saplens mRNA for KIAA0289 gene, partial cds	Homo sapiens GRB2-related adaptor protein (GINAP) minutes	be49405.y3 NIH_MIGC_10 Hamo saprens curva drare invacezcococy o curing	Homo sepiens similar to laminin receptor 1 (67kD, ribosomal protein SA) (H. sepiens) (LOC63484), mRNA	3A) (H. saplens) (LOC63484), mRNA	AU118985 HEMBA1 Homo sapiens cDNA clone HEMBA1004785 5
le Exon Prob	Top Hit Database Source	Ļ	L7	Ę	NT	NT	EST HUMAN	Z-I	LN.	Į.	L	¥	¥	¥	NT	L	FZ.	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN		2	EST_HUMAN	ΝΤ	NT	EST_HUMAN	۲	· Ā	EST_HUMAN
Sing	Top Hit Acession No.	11433721 NT	7662051 NT	7662051 NT	7662047 NT				AF231920.1	4505316		AF007544.1	6806918 NT	6806918 NT		-	1	3.0E-90 BF516168.1	3.0E-90 BF516168.1	3.0E-90 BE563833.1	2.0E-90 BE537913.1		1 N 84 / LSDG	2.0E-90 AI138213.1	2.0E-90 AB006627.1	5729855 NT	2.0E-90 AW672686.1	11427320 NT	11427320 NT	2.0E-90 AU118985.1
	Most Similar (Top) Hit BLAST E Value	5.0E-90	5.0E-90	5.0E-90	5.0E-90	5.0E-90 A	5.0E-90			4.0E-90	4.0E-90.	4.0E-90 /	4.0E-90	4.0E-90	4.0E-90 D87675.1	4.0E-90 /	4.0E-90	3.0E-90 E	3.0E-90 E	3.0E-90	2.0E-90	2.0E-90	2,0E-90	2.0E-90	2.0E-90	2.0E-90	2.0E-90	0 E-90		
	Expression Signal	9.16	0.54	0.54	2.41	2.08	4.43	1.93	1.93	3.26	8.09	86.0	1.07	1.07	7.65	2.17	2.33	1.08	1.08	17.81	4.71	2.67	2.67	1.7	1.05	7.31	4 86	4 78		
	ORF SEQ ID NO:	38209	36264	36265	37648			25742	25743	26482	27125	28405				30109	30123		33571	37551			26568	20200		L				35706
	Exon SEQ ID NO:	22891	L	1	L	24948	24938	١_		13822	14428	1	1	١.	1	1	17500	L	L	1_	L	13905	13905	16577		L		L	1	22510
	Probe SEQ ID NO:	10344	10407	10402	11731	12591	12841	295	295	1084	1684	2882	3023	3023	4608	4743	4768	7751	7751	11630	802	1150	1150	9000	3020	4853		0600	8808	0986 0986

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- 1		l	T		T	Т	Τ	Τ	Π	Г	Τ	Τ	Γ	Τ	Т	Τ	Г		Γ	Į,	7	F	Ī	V	U	5	Г	T	1		1	1		Б	þ
Single Exon Probes Expressed in Brain	Top Hit Descriptor	AU118985 HEMBA1 Homo sapiens cDNA clone HEMBA1004795 5'	Homo sapiens myosin, heavy polypeptide 4, skeletal muscle (MYH4), mRNA	Homo sapiens amyoid beta (A4) preoursor protein (protease nextn-ll. Alzheimer disease) (APP). mRNA	Homo saplens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo seplens mRNA for T-box transcription factor (TBX20 gene), partial	Home sapiens mRNA for T-box transcription factor (TBX20 gene), partial	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens Kruppel-like factor 7 (ubiquitous) (KLF7), mRNA	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 3	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 3	601159563F2 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3511118 67	Homo sapiens similar to SALL1 (sal (Drosophila)-like (LOC57167), mRNA	Homo saplens chromosome 8 open reading frame 2 (C8ORF2), mRNA	Homo sapiens soluble interleukin 1 receptor accessory protein (IL1RAP) gene, exon 8, alternative exons 9	and complete cds, alternatively spliced	Homo sapiens mRNA for KIAA0633 protein, partial cds	Homo sapiens KIAA0623 gene product (KIAA0623), mRNA	Homo sapiens cytochrome P450, 51 (lanosterol 14-alpha-demethylase) (CYP51), mRNA	Human retina-derived POU-domain factor-1 mRNA, complete cds	Homo sapiens glutamate receptor, lonotropio, N-methyl D-aspartate 2A (GRIN2A) mRNA	Homo saplens solute carrier family 1 (high affinity aspartate/glutamate transporter), member 6 (SLC1A8),	mRNA	Homo sapiens brefeldin A-inhibited guanine nucleotide-exchange protein 2 (BIC2), mRNA	Homo sapiens SNCA isoform (SNCA) gene, complete cds, alternatively spliced	Homo saplens CGI-15 protein (LOC51006), mRNA	Homo sapiens CGI-15 protein (LOC51006), mRNA	yg44d11.r2 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:35477 5'	Human branched chain alpha-keto acid dehydrogenase mRNA, 3' end			HUM000S381 Liver HepG2 cell line. Homo sapiens cDNA clone s381 31	
Jie Exon Proc	Top Hit Database Source	EST_HUMAN	TN	Į.	N	NT	N.	LN	LZ	L	TN	IN	ΝT	EST_HUMAN	LN	NT		NT	NT	NT	LN.	NT	TN				NT			T_HUMAN				EST_HUMAN	
N N	Top Hit Acession No.	E-90 AU118985.1	11024711 NT	4502168 NT	E-90 AF231920.1	E-90 AF231920.1	E-90 AJ237589.1	E-90 AJ237589.1	E-90 AF264750.1	E-90 AF 264750.1	4507828 NT	E-90 AF096154.1	E-90 AF096154.1	1.0E-90 BE379884.1	11420514 NT	6005720 NT			1.0E-80 AB014533.1	11426910 NT	141940 <i>6</i>		6006002 NT		11426758 NT	11422086 NT	1.0E-90 AF163864.1	11422109 NT	11422109 NT	1			1.1		
	Most Similar (Top) Hit BLAST E Value	2.0E-90	2.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.05-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90 /	1.0E-90	1.0E-90	1.0E-90		1.0E-90/	1.0E-90 /	1.0E-90	1.0E-90	1.0E-90 U91934.1	1.0E-90		1.0E-90	1.0E-90	1.0E-90 /	1.0E-90	1.0E-90	1.0E-90 F	1.0E-90 J04474.1	1.0E-90 /	1.0E-90 /	8.0E-91 D12234.1	
	Expression Signal	1.37	2.8	4.55	1.36	1.43	2:32	2:32	13.22	13.22	2.47	5.56	6.58	1.23	3.33	6.46		1.29	2.58	96.0	0.57	99.0	0.64		2.77	3.73	96.0	1.53	1.53	1.5	1.76	1.49	1.49	9	
	ORF SEQ ID NO:	35707	36446	25720	25805	25805	26098	26099	26137	26138		26703	26704		27342	28271		29758	31295	31460	32240	32475	32965		33345	34555		35062	35063	36490	36896	31001	31002	29542	
	Exan SEQ ID NO:	22510	23214	13078	16516	16516	13454	13454	13487	13487	13846	14033	14033	14408	14632	15626		17128	18385	18538	19240	19455	19889		20240	21412	21863	21894	21894	23253	23643	24904	24904	16912	
	Probe SEQ ID NO:	9860	11447	270	385	366	679	629	713	713	1088	1283	1283	1662	1895	2868		4389	929	5746	6473	6973	7204		7571	8720	9133	9215	9215	10557	10967	12580	12580	4172	

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11667 SW. ASPG_FLAME Q47898 N4 (BETA-NLACETYLGLUCOSAMINYL)-L-ASPARAGINASE PRECURSOR qe70f1 x1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1744365 3' similar to contains EST01579 Hippocampus, Stratagene (cat. #936205) Hamo saplens cDNA clone HHCMC60 similar to ST01579 Hippocampus, Stratagene (cat. #938205) Homo sapiens cDNA clone HHCMC80 similar to Homo saplens gamma-aminobutyric acld (GABA) B receptor, 1 (GABBR1), transcript variant 2, mRNA au49f09.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2518121 3' similar to zi90b04.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:448015 3 Homo sapiens lysophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds Homo saplens lysophosphatidic acid acytransferase-delta (LPAAT-delta) mRNA, complete cds CBP90 mRNA, partial cds Homo sepiens solute carrier family 4, anion exchanger, member 3 (SLC4A3), mRNA Homo sepiens solute carrier family 4, anion exchanger, member 3 (SLC4A3), mRNA 601901624F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4130933 5 10mo sapiens chromosome 22 open reading frame 5 (C22ORF5), mRNA Homo saplens chromosome 22 open reading frame 5 (C22ORF5), mRNA Homo sapiens cyclin-D binding Myb-like protein mRNA, complete cds AU143539 Y79AA1 Homo sepiens cDNA clone Y79AA1002087 5 Homo sapiens epididymal secretory protein (19.5kD) (HE1), mRNA AU143539 Y79AA1 Homo saplens cDNA clone Y79AA1002087 5' Top Hit Descriptor Homo sapiens makorin, ring finger protein, 1 (MKRN1), mRNA AV649878 GLC Homo sapiens cDNA clone GLCBYF08 3' Rattus norvegicus brain specific cortactin-binding protein AV649878 GLC Homo sapiens cDNA clone GLCBYF08 Homo sapiens cyclin-dependent kinase 6 (CDK6) mRNA Homo sapiens mRNA for KIAA1278 protein, partial cds Homo sapiens mRNA for KIAA1278 protein, partial cds CM-BT043-090299-075 BT043 Homo sepiens cDNA Hamo seplens chramasame 21 segment HS210084 Homo sapiens chromosome 21 segment HS21C083 Human Ku (p70/p80) subunit mRNA, complete cds MIR.b2 MIR MIR repetitive elemen Retrovirus-related gag polyprotein Retrovirus-related gag polyproteln EST HUMAN EST HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN NT EST_HUMAN Top Hit Database Source EST_HUMAN EST_HUMAN Þ 뉟 눋 눋 11497611 NT 7110634 11430193 11430193 4502740 Top Hit Acession 11434964 3.0E-91 AB033104.1 3.0E-91 AB033104.1 3.0E-91 AF084530.1 5.0E-91 AA702794.1 5.0E-91 AU143539.1 5.0E-91 AU143539.1 AV649878.1 AV649878.1 AF053768.1 3.0E-91 AL163283.2 ġ 7.0E-91 A1904151.1 BF314682.1 5.0E-91 AI193566.1 4.0E-91 AF156776.1 4.0E-91 AF156776.1 4.0E-91 AL163284.2 AI879995.1 4.0E-91 M77994.1 M30938.1 5.0E-91 7.0E-91 5.0E-91 (Top) Hit BLAST E Value 5.0E-91 5.0E-91 4.0E-91 3.0E-91 3.0E-91 3.0E-91 3.0E-91 1.93 9. 0.68 .25 1.52 1.69 1.69 1.96 3.07 3.39 5.82 0.97 1.62 3.79 Expression Signal 26859 34036 36062 28877 29840 29841 ORF SEQ 33931 34495 34496 28612 31082 28746 28863 28864 29159 29915 28611 32256 36775 ÖΝΩ 17215 14174 20899 19285 SEQ ID 24919 15960 24596 2886 16223 17215 20799 21350 15960 14360 16094 16520 19255 14360 16211 16211 Exa 18987 ö SEQ ID 8205 3467 4480 10198 6519 8105 8658 8658 12612 12094 **4**88 3197 1613 1613 3334 3455 3455 3768 **8**88 4757 4757 10848 5600 4551 6212

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Trient 2, mRNA thetia 1 (GSTT1)	10557
oript variant 2, mRN, 1 erase theta 1 (GST) 2735280 3°	
Top Hit Descriptor Top Hit Descriptor Source Source Source Source Top Hit Descriptor Source Top Hit Descriptor Source Source Top Hit Descriptor NT Human L-type celclum channel beta-1 subunit (CACNLB1) gens, axons 10 and 11 Human L-type celclum channel beta-1 subunit (CACNLB1) gens, axons 10 and 11 NT Human mRNA for very low density lipoprotein recaptor, complete cds worms 10 and 11 NT Human mRNA for very low density lipoprotein perspective complete cds NT Homo septems mRNA for KIAA1080 protein, partial cds NT Homo septems mRNA for KIAA1080 protein, partial cds NT Homo septems mRNA for KIAA1080 protein, partial cds NT Homo septems mRNA for KIAA1080 protein, partial cds NT Homo septems mRNA for KIAA1080 protein, partial cds NT Homo septems predicting togeths wan 6 Homo septems between the complete cds and the completes of the completes between the completes between the completes of the completes between the completes between the completes between the completes between the completes protein the completes and the completes between the completes and the completes and the completes and the complete completes and the completes and the completes and the completes and the completes and the completes and the completes and the completes and the completes and the completes the completes and the completes and the completes the complete completes and the c	601273513F1 NIH_MGC_20 Homo sepiens cDNA clone IMAGE:3614667 5
Top Hit Database Source Source Source Source NT NT NT NT NT NT NT NT NT NT NT NT NT	EST_HUMAN
A A A A A A A A A A A A A A A A A A A	32 BE 386363.1 E
Acet Similar (Top) Hit Top Hit BLAST E Value 3.0E-91 (186959. 3.0E-91 (186959. 3.0E-91 (186959. 3.0E-91 (186959. 3.0E-91 (186959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-91 (196959. 3.0E-92 (196	8.0E-92 BI
Expression Signal 5.82 4.97 4.97 7.17 7.17 7.17 7.15 0.92 2.25 2.26 2.26 2.26 2.26 2.26 2.26 2.2	828
ORF SEQ ID NO: 33267 33307 34508 35023 35023 36725 37105 37105 37105 377	25/28
Exan SEQ ID NO: NO: 20208 2327 23486 23486 23486 23486 23486 23486 12837 17898 17898 12877 19577	99051
Probe SEQ ID NO:: NO:: NO:: NO:: NO:: NO:: NO:: NO:	8

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	Top Hit Descriptor	au83h08.x1 Schnelder fetal brain 00004 Homo sapiens cDNA clone IMAGE:2782911 3: similar to TR:090302 O60302 KIAA0555 PROTEIN.;contains element MER22 repetitive element;	Homo sapiens mixiva for nicky to be process, process of the phosphatase PYVE-DSP2 mRNA, complete to the containing dual specificity protein phosphatase PYVE-DSP2 mRNA, complete	Polito sapirario i i ve doninario della contra cont	Homo sapiens MCP-4 gene Homo sapiens MCP-4 gene	Homo sapiens DINA polyniciase bear demayer commission	Home sapiens Aliv-1 protein (mo19) gene, exch 11	Human lans membrane protein (mp19) gene, exch 11	Human series transcribition termination factor, RNA polymerase II (TTF2), mRNA	Home conjone mRNA for KIAA0611 protein, partial cds	TOTAL SALVAN OF MARAIL PROBLEM	HOMO SABGETS TILLY IN WICHE PLOCES	Homo sapiens nuclear milogen- and stress-activated protein kinase-1 (MSK1) mRNA, complete cds	Homo saplens dihydrolipoamide S-succinylitransterase (EZ coniponent of Zaxo-gramma compress)	mRNA	Home septens Dive, with case 1 ages,	Trumman Will William according to the Management of the Management	Homo septents mand for KIAA0758 protein, partial cds	Home seniens cycholasmic Seprese truncated Isoform mRNA, complete cds	Home saplens B-cell CLL/Ivmphoma 7b (BCL78) mRNA	Hano sapiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Homo sepiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Hamo sapiens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo saplens NRAS-related gene (D1S155E), mRNA	Homo sapiens DNA, MHC class I region, 7.1 ancestral haplotype	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo saniens T-cell Amphoma invasion and metastasis 1 (TIAM1) mRNA	N-CAM=145 kda neural cell adhesion molecule [human, small cell lung cancer cell line OS2-R, mRNA, 2960]	nt] N-CAM=145 kda neural cell adhesion molecule [human, small cell lung cancer cell line OS2-R, mRNA, 29464] H-CAM=145 kda neural cell adhesion molecule [human, small cell lung cancer cell line OS2-R, mRNA, 79464]	րդ
Sirigle Exoli Flore	Top Hit Database Source	EST HUMAN	뉟	Ŋ	NT	Į.	Ł	IN.	N	į.	Ł	LN	N _T		占	L	Z	Ę!	Z	Z	2 2	1	12	F 1	12 2	N N	1 to		N L	L'N
Silig	Top Hit Acession No.		AB046820.1	AF264717.1	AJ000979.1	AF179428.1	11416961 NT	L04193.1	L04193.1	11426569 NT	AB014511.1	Y13829.1	AF074393.1		4503340 NT	AB031007.1	M60676.1	AB018301.1	AB018301.1	AF007822.1	4502364 N	303137	5051570 NINI	AF 107 7 00. 1	TIM PERSONS	7.0E-92 ABUSTUO7.1		1 N 000 1004	7.0E-92 S71824.1	7.0E-92 S71824.1
	Most Similar (Top) Hit BLAST E Value	8.0E-92 A	8.0E-92 A	8.0E-92	8.0E-92	8.0E-92		8.0E-92 L	8.0E-92 L	8.0E-92	8.0E-92	8.0E-92	8.0E-92		8.0E-92	7.0E-92 /	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.05-92	7.0E-92	7.0E-92	7.0E-92	
	Expression Signal	0.96	0.65	0.97	1.31	0.92	0.55	3.91	3.91	0.58	2.82	1.18	3.2		1.61	1.62	1.01	0.87	0.87	1.34	1.99	227	2.27	1.46			1.06	1.06	2.59	2.59
	ORF SEQ ID NO:	30374	30771	30918	32217	32222		34152	34153	34253	34801	35771			37329	25466	25525		25681					28020				28751	29912	28913
	Exan SEQ ID NO:	17,68	18113	18210	19218	18223	20685	21017	21017	21115	1	l		JACS	24024	L	L	<u> L</u>	15538	L.	14006			15282		15462	17877	17877	17282	17282
	Probe SEQ ID NO:	5041	9308	2.7	845	6455	2862	8324	8324	8422	8960	9925		10/0/	11333	23	49	230	230	577	1257	2184	2184	2568	2728	2757	3340	3340	4547	4547

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Top Hit Descriptor	Homo saplens chromosome 21 segment HS21C081	zw66d12.rf Soares_testis_NHT Homo sapiens cDNA clone IMAGE:781175 5'	801283012F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3605018 6'	601501242F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE:3802939 57	EST91020 Synovial sarcoma Horno saplens cDNA 5' end similar to similar to ribosomal protein S13	Human mRNA for alpha-actinin	Human mRNA for alpha-actinin	Homo sapiens activin A receptor, type IIB (ACVR2B) mRNA	Horno sapiens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	Homo sapiens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	601118337F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3028304 5'	601118337F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3028304 5'	mrg=mas-related [human, Genomic, 2416 nt]	wk27d07.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2413549 3' similar to TR:Q12844 Q12844 BREAKPOINT CLUSTER REGION PROTEIN;	wk27d07x1 NCI_CGAP_Bm25 Home saplens cDNA clone IMAGE:2413549 3' similer to TR:Q12844 Q12844 BREAKPOINT CLUSTER REGION PROTEIN ;	Homo sapiens syndecan 4 (amphiglycan, ryudocan) (SDC4) mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens stress-Induced-phosphoprotein 1 (Hsp70/Hsp90-organizing protein) (STIP1), mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	DKFZp434C0414_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C0414 5'	Homo sapiens P-glycoprotein (mdr1) mRNA, complete cds	Homo sapiens Integrin, alpha L (antigen CD11A (p180), lymphocyte function-associated antigen 1; alpha polypeptide) (ITGAL) mRNA	Homo sapiens mRNA for KIAA1068 protein, partial cds	Human NPY Y1-like receptor pseudogene mRNA, complete cds	Human NPY Y1-like receptor pseudogene mRNA, complete ods	hd02h02.xI Soares_NFL_T_GBC_S1 Home sepiens cDNA clone IMAGE:2908371 3' similar to TR:002711 002711 PRO-POL-DUTPASE POLYPROTEIN ;	Homo saplens thyrold stimulating hormone receptor (TSHR), mRNA
Top Hit Datebase Source	NT	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	LZ.	F.	TN	TN	NT	EST_HUMAN	EST_HUMAN	Z	EST HUMAN	EST HUMAN	Į.	NT	LN	LΝ	LZ LZ	N.	EST_HUMAN	TN	LZ	F	Ž	Ł	174.1 EST_HUMAN	TN
Top Hit Acession No.	2 AL163281.2	7.0E-92 AA446208.1			_	(15804.1	(15804.1	4501898 NT	11422946 NT	2.0E-92 11422946 NT	3E299190.1	2.0E-92 BE299190.1	١.	2.0E-92 AI818119.1	2.0E-92 AIB18119.1	4506860 NT	6912457 NT		2.0E-92 AF231919.1	5803180 NT	A10976.1	2.0E-92 AL040437.1	2.0E-92 AF016535.1	4504756INT	AB028991.1	U87780.1	167780.1	2.0E-92 AW340174.1	11434900
Most Similar (Top) Hit BLAST E Value	7.0E-92	7.0E-92	5.0E-92	3.0E-92	3.0E-92	3.0E-92 X15804.1	3.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92/	2.0E-92 /	2.0E-92	2.0E-92	2.0E-92 /	2.0E-92/	2.0E-92	2 0E-92 /	2.0E-92/L	2.0E-92 U67780.1	2.0E-92	2.0E-92
Expression Signal	0.98	6.05	1.28	1.6	2.6	2.72	2.72	1.68	3.57	3.57	1.33	1.33	2.22	2.38	2.38	5.58	19.2	2.61	2.61	5.67	1.23	2.51	0.64	9.0	3.03	0.81	0.78	1.69	4.93
ORF SEQ ID NO:	30280	30504		28215	31506		1				26162	26163		27378	27379	27504	28113	28992	28993	29068	29626		31383		32285			34594	}
Exon SEQ ID NO:	17871	17989	14328	15473	18577	23355	23365	12852	12986	12986	13506	13506	14462	14665	14665	14776	15374	16353	16353	16427	16997	17664	18468	18984	19282	20045	20045	21446	23350
Probe SEQ ID NO:	4844	5180	1582	2768	6788	10664	10664	24	174	174	732	732	1709	1929	1929	2041	2684	3600	3600	3674	4258	4936	5673	8200	6617	7364	7387	8754	10659

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Top Hit Descriptor	Homo sapiens zinc finger protein 198 (ZNF198), mRNA	Simple Spanic Foliated (Crosspanic) and Crosspanic Cros	Homo sapiens minna for Ninna for the Color of the Color o	Homo sapiens carcineurin piruling protein ((120000), mm	Sares placenta Nozhir nomo sapiaris como sapiaris como ma como como	y80e08.r1 Soares placenta Nb2HP Homo sapiens cunne invade 1400/4 3	Homo saplens ribosomal protein, large, F1 (RFLF1) mixton	HTM1-288F H I M1 Homo sapiens conva	tgotboz.x1 NCI_CGAP_CLL1 Hand seperal SCINA Carlo missociative element; contains element Q16825 PROTEIN-TYROSINE PHOSPHATASE D1; contains Alu repetitive element; contains element MER17 repetitive element;	reotbo2x1 NCI CGAP CLL1 Homo sapiens cDNA clone IMAGE:2107467 3' similar to SW:PTNF_HUMAN	Q16925 PROTEIN-TYROSINE PHOSPHATASE D1 ;contains Alu repetitive element contains element	MER17 repetitive element;	AU121681 MAMMA1 Homo sapiens curve did is invaring to the control of the similar to ribosomal actives and similar to ribosomal	protein L29	is calcium channed apria in squaint (co.co.).	601281867F1 NIH MGC 44 Home sapiens cDNA clone IMAGE:3603832 5		UI-H-Bl0-aah-h-06-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:27093713'	UI-H-BIO-eah-h-06-0-UI.s1 NCI CGAP Sub1 Homo sapiens cDNA clone IMAGE:2709371 3	601460521F1 NIH_MGC_66 Hano sapiens cDNA clane IMAGE:3863908 5	Homo sapiens chromosome 21 unknown mRNA	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1), mKNA	Homo sapiens mRNA for KIAA1267 protein, partial cos	Homo sapiens PTH-responsive osteosarcoma B1 protein (D1) minnin, culturae cus	Homo sapiens mRNA for KIAAU011 protein, parual cus		WOUGOUS X1 NCT CLOSA THING Saprants COT XX COT CLOSA THE COT XX COT CLOSA THE CLOSA THE COT CLOSA TH	odene, complete cds	
Top Hit Database Source					HUMAN	T HUMAN		EST HUMAN HTM1-288F	tgo1502.x1 N Q16825 PRO EST HIMAN MER17 repet	T		\neg	EST HUMAN AU121681 M	EST_HUMAN protein L29		HIMAN		T HI IMAN	Т	HIMAN	Т			NT Homo sapler		Т	HUMAN		יחתוומו פתו
Top Hit Acessian No.	11434759 NT	5803103	AB02901	6912457	R78078.1	R78078.1	450668	BE439625.1	0000	Alseusso.1		AI380356.1	AU121681.1	AA316723.1 ES		AF 223391.1	44.440526	1141624	AW014042.1	AVV U 4042.	AF231919.1	11526176	AB033093.1	AF095771.1	AB014511.1	AI674184.1	AI674184.1	X04201.1	M22878.1
Most Similar (Top) Hit BLAST E Value	2.0E-92	2.0E-92	2.0E-92	2.0E-92	1.0E-92	1.0E-92	1.0E-92	1.0E-92		1.05-92		1.0E-92	9.0E-93	9.0E-93	200	20-20-8 20-20-8	9.05-63	9.05-93	8.05-93	8.0E-93	20-10-50 7 0F-93			\			5.0E-93		5.0E-93
Expression Signal	1.48	2.54	2.69	2.51	1.13	1.13	8.83	1.29		3.82		3.82	8	5.48		1.45	1.11			0.58							7.28		1.01
ORF SEQ ID NO:	36859	36906	31048	28113	27294	27295	27525	١		34910		34911				ı	28995				32203	1				L		28643	31425
Exen SEQ ID NO:	23610	23653	24809	15374				1		21751		21751	1	l _	1]			L	13048	L	L		L		1	18503
Probe SEQ ID NO:	10930	10978	12439	12697	CVa1	18.42	2000	8145		9062		9082	2023	2035		2653	3602	11645	6351	6351	6496	239	200	930	1350	1386	1386	322	5710

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Homo sapiens wbscr1 (WBSCR1) and wbscr5 (WBSCR5) genes, complete cds, alternatively spliced and 250e09.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:795688 3' similar to SW:CLPA_RAT P37397 CALPONIN, ACIDIC ISOFORM; Homo sapiens protein phosphatase-1 regulatory subunit 7 (PPP1R7) gene, exon 11, complete cds and yb94c12.r1 Stratogene liver (#837224) Homo sapiens cDNA done IMAGE:78838 5' similar to similar to SP:A44391 A44391 SERUM RESPONSE ELEMENT-BINDING PROTEIN SRE-ZBP - HUMAN , Homo saplens secretory pathway component Sec31B-1 mRNA, alternatively spliced, complete cds Homo sapiens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA Homo septens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA Homo septens hypothetical protein FL/20731 (FL/20731), mRNA Homo sapiens dystrophin (DMD) gene, deletion breakpoints 1-3 in Intron 5 Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds Homo sapiens chromosome 21 segment HS21C101 Homo saplens discs, large (Drosophila) homolog 2 (chapsyn-110) (DLG2) mRNA Homo saplens discs, large (Drosophila) homolog 2 (chapsyn-110) (DLG2) mRNA h.29g03.x1 NCI_CGAP_Brn25 Homo saptens cDNA clone IMAGE.2169076 3' h.29g03.x1 NCI_CGAP_Brn25 Homo saptens cDNA clone IMAGE.2169076 3' AV692051 GKC Homo sapiens cDNA clone GKCDRF07 5' 602246554F1 NIH_MGC_62 Homo sapiens oDNA clone IMAGE:4332036 5' 602246564F1 NIH_MGC_62 Horno sapiens cDNA clone IMAGE:4332036 5' Horno sapiens tensin mRNA, complete ods Homo sapiens TAR (HIV) RNA-binding protein 1 (TARBP1) mRNA Top Hit Descriptor Homo sapiens gamma-glutamytransferase 1 (GGT1), mRNA Homo saplens Interferon gamma receptor 1 (IFNGR1) mRNA Homo seplens WSB1 protein (WSB1) mRNA, complete cds Homo sapiens gamma-glutamytransferase 1 (GGT1), mRNA Homo sapiens interferon gamma receptor 1 (IFNGR1) mRNA eplication factor C subunit 2 (RFC2) gene, complete cds Homo sapiens Interleukin 18 receptor 1 (IL18R1) mRNA Homo sapiens tumor entigen SLP-8p (HCC8), mRNA Homo saplens nucleobindin 2 (NUCB2), mRNA Single Exon Probes Expressed in Brain alternatively spliced product Top Hit Database HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN Source EST_HUMAN EST_HUMAN EST ż 눋 4557526 NT ż 4557526 NT 11439599 NT 7657454 NT 7657454 NT 11417877 NT 4557879 NT 4557879 N Top Hit Acession 5032156 11417877 8923658 5.0E-93 AF045555.1 AF067136.1 4F069313.2 ġ 5.0E-93 AF274863.1 AA459933.1 4.0E-93 AV692051.1 3.0E-93 BF690630.1 3.0E-93 BF690630.1 3.0E-93 AF225896.1 AL163301.2 4.0E-93 AF047677.1 4.0E-93 AF157476.1 3.0E-83 AI663863.1 3.0E-93 AI553853.1 4.0E-93 | T46864.1 5.0E-93 5.0E-93 5.0E-93 5.0E-93 5.0E-93 4.0E-93 (Top) Hit BLAST E 4.0E-93 4.0E-93 5.0E-93 4.0E-93 4.0E-93 Vost Simila 4.0E-93 Value 1.02 2.15 6.52 Expression 4 1.39 1.53 1.67 1.67 4.3 0.98 1.65 6.17 7.35 7.35 1.31 0.79 0,70 1.67 ORF SEQ 33388 34338 35360 35821 36656 30959 26186 30959 26187 26577 27424 ÖΝΩ 31264 37023 29035 31412 31413 SEQ ID 18799 20280 21195 22176 22358 22618 23415 25052 25052 13218 15118 16308 ö 13914 18354 23748 13527 16396 1291 16396 16951 18491 18491 Probe SEQ ID 6018 8503 8503 9523 7614 9970 12805 88 10727 12343 432 755 3553 ö 432 1970 5557 3643 3643 2397 6697 5697

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	Top Hit Descriptor	Homo sapiens GCN5 (general control of amino-acid synthesis, yeast, homolog Hike 2 (GCN5L2), mRNA	wb02d05.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2304489 3'	Chlorocabus aethiops mRNA for ribosomal protein S4X, complete cds	Chlorocebus aethiops mRNA for ribosomal protein S4X, complete cds	Homo sapiens chromosome 21 segment HS21C085	Homo sepiens chromosome 21 segment HS21C085	Homo sapiens tensin mRNA, complete cds	Human Clk-associated RS cyclophilin CARS-Cyp mRNA, complete cds	601117586F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3358220 5	EST376458 MAGE resequences, MAGH Homo saplens cDNA	Homo sapiens deafness, autosomal dominant 5 (DFNA5), mRNA	QV3-HT0513-290300-126-h04 HT0513 Homo sepiens cDNA	Homo sapiens hypothetical protein (LOC51318), mRNA	HSU74313 Human chromosome 14 Homo sapiens cDNA clone 1-86	UI-HF-BN0-aks-g-09-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA done IMAGE:3078329 5'	AV721846 HTB Homo sapiens cDNA clone HTBAUB04 5'	AV721846 HTB Homo sapiens cDNA clane HTBAUB04 5'	2/29c10.s1 Soares_pregnant_uterus_NbHPU Homo saplens cDNA done IMAGE:503346 3'	Homo sapiens CYP17 gene, 5' end	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	Homo saplens CTR1 pseudogene	Homo sapiens CTR1 pseudogene	Homo sapiens hypothetical protein (DJ328E19.C1.1), mRNA	oy84b08x1 NCI_CGAP_CLL1 Homo seplens cDNA clone IMAGE:1672503 3' similer to TR:Q62384 Q62384 ZINC FINGER PROTEIN.;	Homo saplens DNA for amyloid precursor protein, complete cds	Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD67, mRNA	Homo sapiens hypothetical protein FLJ20291 (FLJ20291), mRNA	Homo sapiens hypothetical protein FLJ20291 (FLJ20291), mRNA	Homo saplens mRNA for KIAA1563 protein, partial cds	Homo sapiens cysteine-rich repeat-containing protein S52 precursor, mRNA, comptete cds	Homo saplens long chain potyunsaturated fatty acid elongation enzyme (HELO1) mRNA, complete cds	
	Top Hit Database Source	IN	EST HUMAN	LΝ	FN	Ę	Ę	ΙN	LΝ	EST_HUMAN	EST_HUMAN	Ę	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	LN	M	M	EST_HUMAN	Ę	Ę	둗	Ę	TN	NT	NT	
	Top Hit Acession No.	11426182 NT	33 AI824829.1	33 AB015610.1	2.0E-93 AB015610.1	2.0E-93 AL163285.2	2.0E-93 AL163285.2	33 AF225896.1		2.0E-93 BE252982.1	2.0E-93 AW964385.1	8153	2.0E-93 BF351469.1	30039	174313.1	2.0E-93 AW 502002.1	1721846.1		2.0E-93 AA126735.1				1.0E-93 AF238997.1	7657016 NT	1		4503872 NT	8923270 NT	8923270 NT		1.0E-93 AF167706.1	1.0E-93 AF231981.1	
	Most Similar (Top) Hit BLAST E Value	3.0E-93	3.0E-93	2.0E-93 /	2.0E-93 /	2.0E-93 /	2.0E-93 /	2.0E-93	2.0E-93 (2.0E-93	2.0E-93	2.0E-93	2.0E-93	2.0E-93	2.0E-93 U74313.1	2.0E-93	2.0E-93 /	2.0E-93/	2.0E-93 /	2.0E-93 L41825.1	2.0E-93 E	1.0E-93 A	1.0E-93 A	1.0E-93	1.0E-93 A	1.0E-93 D87675.1	1.0E-93	1.0E-93	1.0E-93	1.0E-93 A	1.0E-93 A	1.0E-93 A	
	Expression Signal	1.21	4.27	7.51	7.51	9:36	9.48	1.33	1.33	1.66	5.42	99.0	1.04	1.13	0.65	1.1	1.27	1.27	2.64	2.17	10.79	229	2.29	4.25	4.57	8.91	2.9	7.65	7.65	1.5	0.99	1.33	
	ORF SEQ ID NO:	32236	36631	25637	25638				27587	27949		30824		31256	31270		36946	36947				25564	25565	25923	25994	26292	26560	26632	26633	26744	26746	27798	
	SEQ ID NO:	19235	23394	12998	12998	13119	13119	14357	14857	15207	18135	18145	18254	18347	18362	19347	23686	23686	24686	24736	24899	12927	12927	13289	13366	13622	13899	13965	13965	14071	14073	15061	
ſ	Probe SEQ ID NO:	6468	10703	185	185	315	316	1610	2126	2490	5332	5342	5455	5550	5565	6584	11014	11014	12233	12314	12571	5	ē	505	989	852	1144	1215	1215	1322	1324	2337	

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens MHC class 1 region	Novel human gene mapping to chomosome 1	601177686F1 NIH_MGC_17 Hamo saplens cDNA clone IMAGE:3532965 5	601177686F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3532965 5	Homo saplens DNA for amyloid precursor protein, complete cds	Homo sapiens long chain polyunsaturated fatty acid elongation enzyme (HELO1) mRNA, complete cds	Homo sapiens chromosome 21 segment HS210084	Homo sapiens glucocorticold receptor (GRL) gene, intron D, exon 5, and intron E	Homo sapiens glucocorticold receptor (GRL) gene, intron D, exon 5, and intron E	Homo saplens candidate taste receptor T2R14 gene, complete cds	Homo sepiens neurofibromin 1 (neurofibromatosis, von Recklinghausen disease, Watson disease) (NF1)	Homo sapiens KIAA0872 nane product (KIAA0872), mRNA	Homo saplens protein khase C, beta 1 (PRKCB1), mRNA	Human mRNA for NF1 N-Isoform-exon11, complete cds	Homo sapiens mRNA for KIAA1411 protein, partial cds	H.sapiens mRNA for MEMD protein	Homo saplens protein kinase inhibitor gamma (PKIG) mRNA, complete cds	Homo sapiens mRNA for KIAA1485 protein, partial cds	Homo sapiens Trio isoform mRNA, complete cds	Human PreA4 gene for Alzheimer's disease A4 amyloid protein precursor (exon 9)	Human PraA4 gene for Alzheimer's disease A4 amyloid protein precursor (exon 9)	Novel human gene mapping to chomosome 13, similar to rat RhoGAP	Homo sapiens ryanodine receptor 3 (RYR3), mRNA	Homo sapiens GGT1 gene, exon 1	Homo saplens glutathione S-transferase theta 2 (GSTT2), mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, complete cds	Homo sapiens chromosome 21 segment HS210009	omplete cds			zg87g08.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:408594 3'
gle Exon Pro	Top Hit Database Source	N	TN		EST_HUMAN	ΙN	FZ	L'N	L	ΙΝ	LN.	FN	Į.	N-	LN	٦	LN	Į,	ΤN	۲N	TN	NT	LN	NT	IN	Ę		N.	ΙN	Į.	NT	LN L	EST_HUMAN
ଞ୍ଚ	Top Hit Acession No.	-93 AF055068.1	1.0E-93 AL137200.1	1.0E-93 BE297369.1	:-93 BE297369.1	:-e3 D87675.1	1.0E-93 AF231981.1	1.0E-93 AL163284.2	J78509.1	:-93 U78509.1	93 AF227138.1	TN C0277397	7682241 NT	11431590 NT	-93 D42072.1	-93 AB037832.1	-93 Y10183.1	-93 AF182032.1	-93 AB040918.1	-93 AF091395.1	-93 X13474.1	.03 X13474.1	-93 AL049801.1	11433646 NT	1.0E-93 AJ230125.1	11417856 NT		:-93 AF240786.1	8.0E-94 AL163209.2	\F142482.1	5.0E-94 AB014512.1	5.0E-94 AB014512.1	AA722434.1
	Most Similar (Top) Hit BLAST E Value	1.0E-93 /	1.0E-93 /	1.0E-93	1.0E-93	1.0E-93 [1.0E-93	1.0E-93 /	1.0E-93 U78509.1	1.0E-93 (1.0E-93 /	4 OF 03	4 OF 93	1.0E-93	1.0E-93 [1.0E-93/	1.0E-93	1.0E-93 /	1.0E-93 /	1.0E-93/	1.0E-93 >	1.0E-93	1.0E-93 /	1.0E-93	1.0E-93 /	1.0E-93		1.0E-93 /	8.0E-94 /	6.0E-94	5.0E-94	5.0E-94	5.0E-94 /
	Expression Signal	1.7	96.0	1.47	1.47	7.48	1.27	2.6	2.38	2.38	96.0	5 5	1 4	2.01	3.49	2.54	1.2	1.38	1.79	1.26	8.29	8.29	62.0	0.62	1.84	. 284		1.49	1.03	1.63	4.23	4.23	4.22
	ORF SEQ ID NO:	27924		26689	26690	28349		29783	31173	31174	31389	24.548	31850	32650	32907	33984	34269	34382	33542	33548	35319	35320	35467	35910						29333	30747	30748	31698
	Exon SEQ ID NO:	15184		l		15700	16973	17132			18472		1	1	l	20862	21133	l)		22141		22278	22693	24846	24896		25391	23138	16694		1	18739
	Probe SEQ ID NO:	2486	2511	2826	2825	2934	3210	4396	6479	5479	5678	8008	8404	9694	7150	8168	8441	8647	8352	8358	9488	9488	9626	10045	12487	12566		12739	10492	3944	5283	6283	5957

Page 411 of 536 Table 4 Single Exon Probes Expressed in Brain

	П			Т	٦	Т	┪	Ţ	8	\top	7	7,	T	T	Γ	F		7	7	7	"	3	٦	ű,	71	7	П	Щ	1	1	تمد لا
Top Hit Descriptor Top Hit Descriptor	od83d05.s1 Soares_total_fetus_NbZrtrs_9w rightnesseptens conv. concentrations to the concentration of the conference of	002042 1037 1 NOT COMPANY Splen 1NFLS Homo saplens cDNA clone IMAGE:116239 3	yaseputist States locality of processing and the Kitabon 27 professions and the Kitabon 27 pr	Homo saplens cleavage and polyadenylation specific factor 1, 160kD subunit (CPSF1), mRNA	norm capture of the second nordeln L27 mRNA, complete cds	Home sariens motern phosphatase 1, regulatory subunit 10 (PPP1R10) mRNA	mach 2 x1 Spares NFL T GBC S1 Homo sepiens cDNA clone IMAGE:2701679 3'	mag412 x1 Sparas NFL T GBC S1 Homo sapiens cDNA clone IMAGE:2701679 3'	Invitio x1 NCI CGAP Bm52 Homo saplens cDNA clone IMAGE:2259403 3' similar to TR:Q15265 Q15265	PROTEIN TYROSINE PHOSPHATASE;	Home sapiens soluts carrier family 22 (organic cation transporter), member 1-like (SLC22A1L), mRNA	Homo saplens solute carrier family 22 (organic cation transporter), member 1-like (SLC22A1L), mRNA	Homo sapiens huntingtin (HU) gene, excn 3/	Homo saplens hypothetical protein FLU12450 (FLU12450), mixto	Homo sapiens ASHZL gene, complete ous, silings to complete ous, silings	Home sapiens complement compared to the state of the sapients	Home saptens cysteme-rich repeat-containing process of process of mRNA, complete cds	Home capters cystemers and the protect of FP300 mRNA	Homo sapiens E. A. birtung process root, see a contract the sector (HLE), mRNA	Tronio saniens hepatic leukemia factor (HLP), mRNA	TWASAMB 11 Soeres total fetus Nb2HF8 9w Homo sapiens cDNA clone IMAGE:774782 5	Homo saplens zinc finger protein 277 (ZNF277), mRNA	Home saniens mRNA for MEGF2, partial eds	Homo sapiens chromosome 21 open reading frame 18 (C210RF18), mRNA	Homo sapiens neuronal cell adhesion molecule (NRCAM) mRNA	Homo saplens protocacherin alpha 13 (PCDH-alpha13) mRNA, complete cds	Homo sanians mRNA for KIAA0679 protein, partial cds	Homo saniens chrodenir-1L mRNA, complete cds	Homo sapiens axonal transport of synaptic vesicles (ATSV) mRNA	Human cbl-b truncated form 1 lacking leucine zipper mRNA, complete cds	
Top Hit Database Source	EST HUMAN		EST HUMAN		Z	ž.	- N	TOTAL TOTAL	מאוסני ופש	EST HUMAN	INT	NT.	LΝ	ZINT	N	NT NT	Z	<u></u>	NT NT	IN C	NAME OF THE PERSON AND THE PERSON AN	-1	110	L L	NIN C	1112	Z I	2 2	- P	12	
Top Hit Acession No.	A1015800.1	BF529115.1	T89398.1	D25217.2	9558724 NI	L05094.1	4506008 N	AW197851.1	AW19/851.1	AI591312.1	11440670 NT	11440670 NT	L27386.1	11545792 NT	AB022785.1	4502506 NT	AF167706.1	4 AF167708.1	4557556 NT	11427779 NT	1142///BINI	3.0E-94 AA4648U5.1	070651.1	AB0115	TIN 82282611		3.0E-94 AF152309.1	3.0E-94 AB014579.1	AF08/94	4/5/8Z1 N1	3.0E-94 U.20/ 11.1
Most Stmilar (Top) Hit BLAST E Value	5.0E-94			5.0E-94	5.0E-94	4.0E-94 L	4.0E-94	4.0E-94 /	4.0E-94	4.0E-94	4.0E-94	4.0E-94	-	4.0E-94	3.0E-94	3.0E-94	3.0E-94	3.0E-9	3.0E-94	3.0E-94											
Expression Signal	1.29	1.11	66.6	1.5	1.86	2.64	0.92	1.38	1.38	2.87	1.82	1.82	1 18	1.5	1.44	0.88			2.61					1.33							1:94
ORF SEQ ID NO:	32693	34371	30619					29053		30038	32143	}		36420		L			Ŀ		27535				32125		2 33923	8 34322			4 37596
Exen SEQ ID NO:	19648	21229	L	1_	25021	L	15372	16414	16414	17403	1	1	1	L	L	┸	L	L	1.	L	3 14805			18839	19130	6 20359	8 20792	8 21178	22145		9 24274
Probe SEQ ID NO:	6910	8537	12209	12756	12761	1834	2662	3661	3661	4669	6378	3 8	2/20	7100	102	3	1733	1733	1765	2073	2073	4166	5595	6029	6360	7696	8098	848	9492	11043	11679

Page 412 of 536 Table 4 Single Exon Probes Expressed in Brain

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Top Hit Descriptor	wi30h11.x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:2391813 3'	wi30h11 x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:2391813 3'	601175762F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3531038 5	601111696F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3352859 6	601111898F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3352559 5	Homo sapiens hypothetical protein (FLJ20748), mRNA	Escherichia coli K-12 MG1655 section 159 of 400 of the complete genome	DKFZp434G0314_11 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434G0314 5	y87f02.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:45053 6'	AV725992 HTC Homo sapiens cDNA clone HTCBEF05 5'	Homo sapiens chromosome 21 segment HS21C004	Homo sapiens chromosome 21 segment HS21C004	Hamo sapiens paired box gene 5 (B-cell lineage specific activator protein) (PAX5), mRNA	601468748F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3872099 6'	Homo sapiens IL-1 receptor antagonist IL-1Ra (IL-1RN) gene, alternatively spliced forms, complete cds	ap22e02.x1 Schiller oligodendroglioma Homo saptens cDNA clone IMAGE:1966122 3' similar to TR:Q62845 Decay 45 NEURAL OF LADHESION PROTEIN BIG-2 PRECURSOR	Homo sapiens KIAA0164 gene product (KIAA0164), mRNA	RAN175767F1 NIH MGC 17 Homo sapiens dDNA clone IMAGE:3631038 6	601175762F1 NIH MGC 17 Homo saplens cDNA clone IMAGE:3631038 5'	Homo saplens TNF-alpha stimulated ABC protein (ABC50) mRNA, complete cds	Homo saplens KIAA0255 gene product (KIAA0255), mRNA	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA	M.musculus glyT1 gene (exans 1c and 2)	M.musculus glyT1 gene (exons 1c and 2)	Homo sapiens progressive ankylosis-like protein (ANK) mRNA, complete cds	we09e04.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2340606 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	we09e04.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2340606 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN);	Homo sapiens KIAA0193 gene product (KIAA0193), mRNA	Homo saplens proteasome (prosome, macropain) 26S subunit, non-ATPase, 11 (PSMD11), mRNA	Homo sapiens proteasome (prosome, macropain) 26S subunit, non-ATPase, 11 (PSMD11), mRNA	
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	LN	EST_HUMAN	F	NAMI H TAT			EST HIMAN	L	_	FN.	NT	NT	M	EST HUMAN	EST HUMAN	NT	LN	N.	
Top Hit Acession No.	AI910393.1	AJ910393.1	BE295714.1	BE253433.1	BE253433.1	9506692	AE000269.1	AL040518.1	H08270.1	AV725992.1	AL163204.2	AL163204.2	11428710 NT	BE780478.1	U65590.1	A1979244 4	11418871	DE-2087444	BE 208714.1	AF027302.1	7662027	7662027	X82569.1	X82569.1	AF274753.1	AI700998.1	AI700998.1	11419376 NT		11428529 NT	
Most Similar (Top) Hit BLAST E Value		2.0E-94 /	1.0E-94	1.0E-94	1.0E-94	1.0E-94	1.0E-94 /	1.0E-94 /	1.0E-94	1 0E-94 /	1.0E-94	1.0E-94	1.0E-94	1.0E-94	1.0E-94	10.10	_			9 OF-95		9.0E-95	9.0E-95	9.0E-95	9.0E-95	8.0E-95			8.0E-95	8.0E-95	
Expression Signal	0.7	2.0	1.94	2.07	2.07	1.7	0.64	0.73	0.72	0.58	0.63	0.63	2.76	2.04	3.08		4 72	7,1	1.40	1 83	1.45	1.45	1.33	1.33	1.77	3.18	3.18	0.76	1.76	1.76	
ORF SEQ ID NO:	35501	35502	25601	28492	28493			31922	31933	32190	33836	33837	35054	35533	36929	21010			20001					30782	33974						
Exon SEQ ID NO:	22305	22305	12959	15851	16851	17065	18763	18950	18959	19194	20707	20707	21886	22339	23673	9	24245	242	12859	14208	15918	15916	18123	18123	20844	ļ	1	1	19828	19828	
Probe SEQ ID NO:	9653	8883	4	3086	3088	4328	5982	6173	6182	6426	8012	8012	9155	2896	11000	3	11200	6	12330	4450	3153	3163	5320	5320	8150	4400	7,00	0708	7141	7141	

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	Top Hit Descriptor	Homo sepiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo sapiens KiAA0255 gene product (KIAA0255), mRNA	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA	Homo saplens proline dehydrogenase (proline oxidase) (PRODH) mRNA	Homo saplens mRNA for KIAA1395 protein, partial cds	Homo saplens early growth response 2 (Krox-20 (Drosophila) homolog) (EGR2), mRNA	Homo sapiens developmental arteries and neural crest EGF-like protein mRNA, complete cds	Homo sapiens HCF-binding transcription factor Zhangfel (ZF), mRNA	zu84b01.s1 Sogres_testis_NHT Homo sapiens CDNA clone IMAGE:744649 3' similar to contains L1.t1 L1	repeauve element;	Homo sapiens DNA for amylad precursor protein, complete cds	Homo saptens DNA for amylad precursor protein, complete cds	Homo sapiens Ly-6-like protein (CD59) mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C046	HTM1-288F HTM1 Hano sapiens cDNA	602071146F1 NCL CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4214147 5	Homo saplens dedicator of cyto-kinesis 1 (DOCK1) mRNA	EST370191 MAGE resequences, MAGE Hamo sapiens cDNA	EST370191 MAGE resequences, MAGE Homo sapiens cDNA	aus3b08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783799 3' similar to TR:060463 060463 TYPE-2 PHOSPHATIDIC ACID PHOSPHOHYDROLASE. [1];	aug3b08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone INAGE:2783799 3' similar to	IN. COURS CONTROL ITTE-2 PROSPINATION AND PROSPINATION CLASS. [1];	Hamber contains Kith ADT62 acres product (KITA 40783) mDNA	INCHES CAPACITES AND AND AND ESTIMATE AND AND AND AND AND AND AND AND AND AND	WRZ7111 States fold live solem UNE S Home seniors CDNA From IMAGE 19448 F.	Homo sariers KIAA0255 cans product (KIAA0255) mRNA	Home serious KIAA0755 cans product (KIAA0755) mm.	Homo sablens fission inhibitor of metalloproteinese 3 (Sorchy findus dystrophy pseudoinflammatro) (TIMDS)	mRNA	601312161F1 NIH_MGC_44 Hamo sapiens cDNA clane IMAGE:3658862 5'	Homo sapiens G protein-coupled receptor 19 (GPR19) mRNA	Homo saplens G profein-coupled receptor 19 (GPR19) mRNA
21 112	Top Hit Database Source	LN	LΝ	Į.	LN LN	LN	TN	ĮN	Į,		ESI HUMAN	Į.	NT	NT	ĹΝ	EST_HUMAN	EST_HUMAN	ΤN	EST_HUMAN	EST_HUMAN	EST_HUMAN		ES HOMAN	1	ECT UI MANI	EST HIMAN	-1.	HN.		NT.	EST_HUMAN	FN	Į,
	Top Hit Acession No.	AF032897.1	11420944 NT	11420944 NT	8.0E-95 5174644 NT	8.0E-95 AB037816.1	9845523 NT	AF1121	10864024 NT	700000	15 AA629056.1	5 D87675.1	D87675.1	7.0E-95 M95708.1	7.0E-95 AL163246.2	4.0E-95 BE439625.1	3.0E-95 BF526041.1	4503354 NT	3.0E-95 AW958121.1	3.0E-95 AW958121.1	3.0E-95 AW157233.1		3.0E-95 AVV15/233.1 ES		DE21244	2 OF OS D83100 1	7662027 NT			4507512 NT		2.0E-95 5453665 NT	5453665 NT
	Most Similar (Top) Hit BLAST E Value	8.0E-95	8.0E-95	8.0E-95	8.0E-95	8.0E-95	8.0E-95	8.0E-95	8.0E-95	L	8.0E-95	7.0E-95	7.0E-95	7.0E-95	7.0E-95	4.0E-95	3.0E-95	3.0E-95	3.0E-95	3.0E-95	3.0E-95	100	3.0E-95	3.05-93	3.05.05	3 OF OF	205-92	20000	207	2.0E-95	2.0E-95	2.0E-95	20E-95
	Expression	1.97	2	2	3.1	2.94	0.8	1.3	1.86	į	12.4	32.81	32.81	5.18	1.3	1.31	1.58	0.83	1.51	1.51	0.55		0.0	8.5	60.	7 40	29.4	224	2	2.51	1.92	2.22	222
	S Ci	33922		_	35604		35994				1			29701	-					33028	34113		25407	١	١			27072	2017	27384		27880	╝
	_ w	20791		ı	l	l			24069					L	1111	!			1	19952	20972	ļ	71802	1	1		14385			14669			_ !
	Probe SEQ ID NO:	8097	9265	9265	9748	9779	10135	10613	11466	1010,	CSC21	987 789	269	4333	4380	9117	5356	5588	7268	7268	8278		8/78	9200	0647	10707	1630	1820	3	1834	1837	2426	2426

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Top Hit Descriptor	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	Homo saplens glycine cleavage system protein H (aminomethyl carrier) (GCSH) mKNA	Homo saplens Usurpin-gamma mRNA, complete cds	Homo sepiens unconventional myosin-15 (LOC51188), mRNA	Homo sapiens unconventional myosin-16 (LOC51168), mRNA	Homo sapiens mRNA for KIAA 1386 protein, partial cds	qm01c02.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1880b46 3' similar to WP:1.23c7.4 CE03705 ;	Homo saplens hypothetical protein (HS322B1A), mRNA	Homo saplens KIAA0187 gene product (KIAA0187), mRNA	Homo sapiens CGI-48 protein (LOC51096), mRNA	Homo sapiens CGI-48 protein (LOC51096), mRNA	Homo sapiens angiotensin I converting enzyme (peptidyl-dipeptidase A) 2 (ACE2), mKNA	Homo sapiens angiotensin I converting enzyme (peptidyl-dipeptidase A) 2 (ACE2), mKNA	Homo sapiens membrane protein, palmItoylated 3 (MAGUK p55 subfamily member 3) (MPP3), mRNA	Human muscle-type phosphofructokinase (PFK-M) gene, exon 7	Homo sapiens transcription factor 2, hepatic; LF-B3; variant hepatic nuclear factor (TCF2), mRNA	Homo sapiens transcription factor 2, hepatic; LF-B3; variant hepatic nuclear factor (1 CF2), mKNA	Homo capiens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds	Homo saplens huntingtin (Huntington disease) (HD), mRNA	Homo sapiens ribophorin II (RPN2), mRNA	Homo saplens KIAA1065 protein (KIAA1065), mRNA	Homo sapiens bone morphogenetic protein receptor, type IA (BMPK1A) mKNA	Homo sapiens KIAA0763 gene product (KIAA0763), mRNA	Homo sapiens KIAA0763 gene product (KIAA0763), mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) and glutathione S-transferase theta 1 (GSTT1)	United contact and an inclinate lines (ADSL) mRNA	Trutho express and specific for the control of the MARCE TAMAN Reliable to	72.23104.11 Sociales ovary unition future in the course ovary unition for	zt23h04.r1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:714007 5' similar to TR:G1067084 G1067084 F55H2.6;	-
Top Hit Database Source	NT	NT	NT	NT	N L	۲N	EST_HUMAN	LN PL	ΙΝ	NT	LΝ	TN	TN	TN	F	TN	LN.	TN	NT	NT.	NT	NT	NT	ΝΤ	Ė		Ž	EST_HUMAN	EST_HUMAN	
Top Hit Acession No.	AF240786.1	4758423 NT	AF015452.1	7705900[NT	7705900 NT	AB037807.1	AI290264.1	7657185 NT	7861979 NT	7705764 NT	7705764 NT	11225608 NT	11225608 NT	11525883 NT	M59724.1	11427182 NT	11427182 NT	AF257737.1	11435773 NT	11421795 NT	11434330 NT	4757853 NT	7662289 NT	7662289 NT	7 00207 011	AF-240780.1	11418164 IN	AA284651.1	AA284651.1	
Most Similar (Top) Hit BLAST E Value	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95 /	2.0E-95		2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95		2.0E-85	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95		2.0E-95		2.0E-95	2.0E-95	1.0E-95		
Expression Signal	3.28	1.85	1.96	3.07	3.07	1.29	0.68	1.32	272	4.21	4.21	1.27	1.27	0.7	5.04	1.16	1.16	2.45	1.82	1.06	0.84	2.46	3.02	3.02			4.66	8.41		
ORF SEQ ID NO:	27925	27973	28564	28958	28957	28998	29122		30308	30884	30885	31319	31320			L	L			34886	_	L	37620				30994	31228		
Exon SEQ ID NO:	15186	ı	1_	1_	I_{-}	1_	l	L		L	L	L			L		L		1_	L	L	L	! _	L	1	- 1	24980	18325	L)
Probe SEQ ID NO:	2468	2517	3155	3662	3552	3605	3732	4328	4978	5392	5392	5811	5611	5651	A0E4	6368	6358	6476	9999	9041	10280	10624	11700	11700		12299	12698	7688	5527	

Page 415 of 536 Table 4 Single Exon Probes Expressed in Brain

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Top Hit Top Hit Descriptor Database Source	RC6-FN0019-290600-011-G11 FN0019 Homo sapiens cDNA	RC6-FN0019-290600-011-G11 FN0019 Homo sapiens cDNA	Jen4237232F1 NIH MGC 72 Homo sapiens cDNA done IMAGE:3922423 5	PMn.1 T0019-090300-002-409 LT0019 Homo sapiens cDNA	Lows carians chromosome 21 unknown mRNA	Trull Sapiett of Schools are dehydrogenase pseudogene 3'end	Tulial gyod many year	Homo sapiens statytransferase 6 (N-acetylacosaminide alpha 2,3-statytransferase) (SIAT6), mRNA	Home sapiens KIAAU/60 gene product (NAA0763), mRNA	nome saperis renoved gotto properties 2 skeletal muscle, adult (MYH2), mRNA	Home salvens mycan, nearly payed and the MANA	Tromo sapirats Michael Series Programmer (KIAA0763), mRNA	Tromo saplens mRNA for KIAA1172 protein, partial cds	Tours series mRNA for KIAA1172 protein, partial cds	Home seniors mRNA for KIAA1172 protein, partial cds	Home saniens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE6A), mRNA	Leaning DNA for monoamine oxidase type A (7) (partial)	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo sanians A kinasa (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sanians mRNA for KIAA0960 protein, partial cds	Homo saniens mRNA for 14-3-3gamma, complete cds	Human type IV collagenase (CLG4B) gene, exon 5	Human tyre IV collegenase (CLC4B) gene, exon 5	Homo seniens KIAA0175 gene product (KIAA0175), mRNA	ura7h12 rf Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:212327 5	Homo saniens chondroitin sulfate proteoglycan 4 (melanoma associated) (CSPG4), mRNA	Homo saniens chromosome 21 segment HS21C048	1073 HT0230 A00500-110-002 HT0230 Homo sapiens cDNA	_	Т	7	7	٦.
Top Hit Database Sœurce	EST HUMAN	COT LI IMAN	TOT TOTAL	TOT TOTAL	ESI HUMAN	Z	Z	NT	L	Z	Z	Į.	Į.	Ž.	Z	Z	2 12	i N		Z	2 1	Z	Z L	- F	12/2	TOT LIMAN	-1	2 1	- N	TOT TOT	TOT TOT	HOLL HOLL	TOT TOTAL	EST HOMAIN
Top Hit Acession No.	1,0000	7 0000	1				73.1	11422642 NT	7662289 NT	7662289 N	8923939 NT	7682289 NT	7662289 NT	5.0E-96 AB032998.1	5.0E-96 AB032998.1	5.0E-96 AB032998.1	11416/0/ N	(60812.1	Ĭ	11424399 N	T1424088	(B023177.1	15024334.1	A68347.1	M68347.1	INIC/BLOO/	168656.1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	4L163248.2	2.0E-96 BE148074.1	2.0E-96 BF369731.1	BF369731.1	2.0E-96 AV689461.1	2.0E-96 AW249440.1
Most Similar (Top) Hit BLAST E	4 OF OF BE37	1.00-30	1.0E-95 BF3/	9.0E-96 BE89/259.1	8.0E-96 A	7.0E-96 AF231920.1	6.0E-96 M268	6.0E-96	6.0E-96	6.0E-96	6.0E-96	6.0E-96	6.0E-96	5.0E-96 A	5.0E-96 A	5.0E-96 A	5.0E-96	5.0E-96 X60812.1	5.0E-96 AF1	5.0E-96	5.0E-96	5.0E-96 AB0	5.0E-96 AB0	5.0E-96 M68	5.0E-96 M68347.1					_				
Expression Signal	- 6	6.4	4.3	1.49	2.77	0.74	20.13	0.74	3.36	3.36	2.05	1.83	1.83	2.74	4.08	4.08	1.43	1.39	1.15								12.32	4.24					5.63	2.81
ORF SEQ ID NO:	11.00	33175	33176	33920		29278	28886	31258		37440		37692		25754		26261			32324							37715			26159	30072		33121	_	
Exon SEQ ID NO:		20091	20091	20788	18223	16639	16232	18349	L	L		1	1_	L		13592	15336	17576	19318	19601	19601	19659	20092	20700	20700	3 24383	16908	13191	13504	3 17440	1 20042	1 20042	١	1
Probe SEQ ID NO:		7414	7414	8094	5424	3889	3476	5550	11534	11534	11584	11769	11769	312	822	822	2624	4846	6553	9684	6684	6923	7415	8005	8005	11793	4168	406	730	4708	7361	7361	8879	12009

Page 416 of 536 Table 4 Single Exon Probes Expresse

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Top Hit Descriptor	Hama contains a survey of the line of the survey of the line of the survey of the line of the survey	Home septens neuronal cell agnesion molecule (NRCAM) mRNA	Himmondon cell ednesion molecule (NRCAM) mRNA	FST287124 MACE	EST387124 MAGE resequences, MAGC Homo sapiens cDNA	Edit catic cincidations and a control of the contro	Home calles superified in nearly chain (sMyHC) mRNA, complete cds	Homo septens dansient redeptor potential channel 6 (TRPC6), mRNA	Home sariens HSDC144 restain 71 100001.	Homo sapiens HSPC144 profein (HSPC144), mRNA	Homo saplens similar to ectonucleotide pyrophosphatase/phosphodiestarase 3 (H. saplens) (LOC83214), mRNA	Homo saniane secretor radium.	Homo senione mBNA for VIA A 1900.	Homo septems mRNA for KIA A 1300 protein, partial cds	Homo sablens perimonal cell adhesion and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contractions and contractions are contracted and contracted and contractions are contracted and con	Tomo sanians perinanal pall pall perinanal pall pall perinanal pall pall perinanal pall pall pall perinanal pall pall pall perinanal pall pall perinanal pall pall perinananal pall perinanananananananananananananananananana	F-HT0417-041000 not not not not not not not not not not	601440317F1 NIH MCC 72 Home collistic CNA	601440317F1 NIH MGC 72 Home emilian child in 11.00 mg and	EST22672 Adipose fissile white II home conice. Only Fi	EST22672 Adipose tissue, white II Home septems con A 5	Human mRNA for alpha-actinin	DKFZp434N0323_r1 434 (synonym: htss3) Homo sapiens cDNA clone DKFZp434Nn324 K/	ZA97e12.s1 Sogres_NHHMPu_S1 Homo septens cDNA clone IMAGE:767758 3 similar to TR:G1304125 U	BCO-BT0813.350000 023 -00 BT2010 II	ARD-HTD241-450600 040 Log Lifes 411	Micro Listan - 10-00-0 10-00-0 10-00-0 Micro Micro Listan - 10-00-0 10-00-0 Micro Micro Listan - 10-00-0 Micro Micro Listan - 10-00-0 Micro Micro Listan - 10-00-0 Micro				Homo sapiens brefeldin A-Inhibited guanine nucleodide-exchange protein 2 (BIG2), mRNA
Top Hit Database Source	ΙN	Į.	L N	EST HIMAN	EST HIMAN		1	L	Į.	FX		Į.	Į.	¥	Ę		T HUMAN	Т	Т	Т	Т	-N	EST_HUMAN	EST HIMAN	T	HIMAN	NAM.		,	Į	
Top Hit Acession No.	4826863	TN SARRES INT	18890 1	1.0E-96 AW955054.1	1.0E-96/AW955054.1	151472.2	6912735INT	6912455 NT	7661803 NT	7661803 NT	11419429 NT	-274863.1	3033116.1	Γ	8883	4826863 NT	141849.1	898012.1	898012.1	Γ	320332.1	804.1	43314.2		T	Τ	T	T			5453572
Most Similar (Top) Hit BLAST E Value	1.0E-98	1.0E-98	1.0E-98 V	1.0E-96/	1.0E-96/	1.0E-96 L	1.0E-98	1.0E-96	1.0E-96	1.0E-98	1.0E-96	1.0E-96 AF	1.0E-96 A	1.0E-96 A	1.0E-96	1.0E-96	6.0E-97 BE	6.0E-97 BE	6.0E-97 BE	8.0E-97 A	6.0E-97 AA320332.1	6.0E-97 X15	5.0E-97 AL	6.0E-97/A/	5.0E-97 BF154912.1	5.0E-97 BE148597.1	5.0E-97 RF148597 1	4.0E-97 BE	4.0E-97 AB030176.1	4.0E-97 AB(4.0E-97
Expression Signal	2.6	2.8	3.38	2.56	2.56	1.06	1.3	0.67	1.24	1.24	20.66	2.09	1.24	1.24	1.97	1.97	2.6	0.69	0.69	0.57	0.57	1.48	1.91	11.73	2.66	1.98	1.98	1.59	1.04	1.04	1.07
ORF SEQ ID NO:	26018	26017	26074	27218	27217	27729	30541			33939	34450	34588	35924	35925	26016	26017		34670	34671	36369	36370	37290	33739	33869	35412	37441	37442	26349	26359	26360	27349
Exon SEQ ID NO:		13386							20805	20805	21308	21441	22707	22707	13386	13386	20131	21524	21524	23143	23143	68857	8008	20736	22227	24135	24135	13685	13695	13895	14840
Probe SEQ ID NO:	808	808	656	1774	1774	2262	6889	6949	8111	8111	8618	8749	10059	10059	2 88	188 88	7457	883	8832	10497	10497	1382		8042	9574	11635	11535	918	928	828	1903

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	Top Hit Descriptor	Lance conjugge explication H (beta-2-divoporotein I) (APOH) mRNA	Hullio Saprieris application () Application (Human N-meu iyi-C-asharan 1000km ii Iiiii N-meu iyi-C-asharan noqulatan subunit 2A (hNR2A) mRNA, complete cds	Furnari I villed yi-Drasparian 1997 - Basia Vitransferase I, long form	Fromo saprens III.NA for GallNAc alpha-2. 6-sialytransferase I, long form	Truling September III TONA ATP-dependent (LIGS), transcript variant alpha, mRNA	Home caries a series fibrais transmembrane conductance regulator, ATP-binding cassette (sub-family C,	member 7) (CFTR), mRNA member 7) (CFTR), mRNA	Homo saprens tanninn, apprais (Schmidt-Ruppin A-2) viral oncogene homolog (SRC), mRNA	House selvers of the property of supplementally IVB polypeptide 1 (CYP4B1), mRNA	Home saprens cyclinated to comment of the saprens o	Homo sapiens minute for KTAAAROA protein, partial cds	Homo sapiens minds for increase from the same (AXL), transcript variant 1, mRNA	Homo sapiens AXL receipe kinese (AXL) transcript variant 1, mRNA	Homo saplens AAL teceptor vironic intractive — //	Home series mRNA for KIAA1290 protein, partial cds	Truits deprete in the second of the second o	Home saplens in NAM to Name see expressed 1 (GTSE1), mRNA	House services of End (AA1172 protein, partial cds		Homo sapiens amyloid beta (A4) precursor protein (protease nedn-II, Atzhelmer disease) (APP), mRNA	Homo sapiens amyoid beta (A4) precursor protein (protease nexin-II, Alzheimer disease) (APP), mRNA	Homo sapiens N-myc (and 3 I A / Itilistator) (1 www),	Human beta-prime adaptul (DAWLE) gales according to a contract and the property mRNA	Home septents personally translation elongation factor 1 alpha 1 (EEF1A1) mRNA	_	\neg	RC1-H70258-211199-011-g05 HT0258 Homo sapiens cDNA	NA clone IMAGE:129134 3'			
Single LAURI FIRE	Top Hit Database Source				LN.	LZ.	LN	Z	N _T	LZ.	LN	L	LZ	Z	2	Z	EN!	N	Z!	Z.	- N	FN	INT	NT NT	<u> </u>	IN .	TNIC	TOT TOT	TOT TOTAL	MAN IL	TIVE TOWNS	TN C	
Rills	Top Hit Acession No.		4557326				Y11339.2	7710125 N	11422155 NT	4557708 NT	11421793 NT	11423233 NT	AB011166.1	AB011166.1	11863122 NT	11863122 NT	AB042557.1	4.0E-97 AB033116.1	4.0E-97 AB033116.1	11418318 NT	3.0E-97 AB032998.1	4502166 NT	4502166 NT	4758813 NT	7.1		4503470 NT	1.0E-97 BE566486.1	1.0E-97 AW379976.1	97 AW379976.1	K1088/	TN 737767 F	
	Most Similar (Top) Hit T BLAST E						4.0E-97 Y	4.0E-97	4.0E-97	4.0E-97	.4.0E-97		4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	4.0E-97	3.0E-97	3.0E-97	3.0E-97	3.0E-97					-1 	1.0	1		1.00
	Expression Signal		0.61	0.95	0.95	6.47	6.47	-	1.05	0.57	2.63	0.75	1.23	1.23	1.88	1.88	3.61	1.62	1,62	7.76	1.14	29.53	29.53				1	2.48					3.44
	ORF SEQ ID NO:		31172	31482	31483	32590	32591	32703	32468	33863	34087							36411			25685	26294	l				L	32091	33534				8 36539
	SEQ ID		18277	18558	١.	1	1	l_	19450	1_	1	1	1	┸	ı	1_	1	L	1_	_	L		1	┸	L		1		L	L	4 . 22316	H	4 23298
	Probe SEQ ID		5478	5785	5765	6725	6725	6921	8909	8036	8258	8518	277	0147	1118	1116	11412	11415	11415	12180	236	854		624	248	325	4729	6333	9344	9344	9664	10604	10604

Page 418 of 536 Table 4 Single Exon Probes Expressed in Brain

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III Dance of the control of the cont	Top Hit Descriptor	Т	7	Home scriptor in the sound protein S18 (RPS16), mRNA	Т	1	House series cat eye syndrome critical region gene 1 (CECR1), mRNA	Home septems CLUN12 gene for claudin-12	Home septents reucyt-trivity synthetese, mitochondrial (KIAA0028), mRNA	Home explains A utiliass (FricA) anchor protein (yotlao) 9 (AKAP9), mRNA	Home serious doct to the contract of the contr	Hingas w BAIA	Home and the sunying A4(751) protein	Upon Septems Succindate-Coa ligase, GDP-forming, alpha subunit (SUCLG1), mRNA	Hours septems mKNA for KIAA1365 protein, partial cds	Homo sapiens 17-beta-hydroxysteroid dehydrogenase IV (HSD17B4) gene, exon 8	Homo sapiens SW I/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 3 (SMARCA3) mRNA	Homo sapiens SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a	member 3 (SMARCA3) mRNA	Homo squiens inosital polyphosphate 1-phosphatase (INPP1) gene, complete cds	nomo sepiens mxnA tor KIAA1005 protein, partial cds	nomo septens mKNA for KIAA1005 protein, partial cds	Home septents mitogen-activated protein kinase kinase Kinase 7 (MAP3K7), mRNA	Homo saplens mRNA for MEGES would be a	Homo sapiens hPAD-calond mRNA for population 11 11 11 11 11 11 11 11 11 11 11 11 11	Homo septiens PMS2146 mRNA control population definite de	Homo sapiens PMS21 16 mRNA pertial cds	Human milochondrial creatine khake (CKMT) con	601507503F1 NIH MGC 71 Homo semicas CONIA ALL MACH CONTROLL CONTRO	Т	Homo sapiens mRNA for KIA A0707 sectors appears cDNA clone is	T	Homo sapiens activator of Subara Unace (Active Control of Control	Homo sapiens activator of S phase kinase ASK1 mBNA	CANALL (VIOL) CORES CORES
	Top Hit Database Source	FOT HIMAN	L L	1	EST HIMAN	• I.	- L	į	FZ.	FZ	LZ	LZ		L Z	H		F	<u> </u>	E	5 5	į.	Į.	Ę	Z	Ę	F	¥	¥	EST HUMAN	ST HUMAN	N	EST HUMAN	1		
	Top Hit Acession No.	-97 AA553761 1	1142R272INT		BE0909	N COURSE	A.1250713 4	9.0E-98 Z661871 NT				X06989 1	13215BO	-98 AB037786 1	Ī		4507070 NT	714 0502027	98 A E141325 2	T		2000	98 AB011541.1		98 AB033768.1				6.0E-98 BE885873.1 E				19210	11419210 NT	
	Most Similar (Top) Hit BLAST E Value	1.0E-97		1.0E-97	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9.0E-98	9 OF-08	2:01	9.0E-98	0 P-0	9.0E-98	9.0E-98	9.05	9 OE-98	9.0E-98/	9.0E-98 /	8.05-98	8.0E-98	8.0E-98	8.0E-98	6.0E-98	3.0E-98 A	3.0E-98 A	3.0E-98 A	3.0E-98	3.0E-98	
	Expression Signal	2.03	14.01	14.01	3.52	1.12	0.71	0.67	9.0	4.79	4.79	6.28	1.5	1.59	0.96		1.14	7.1	0.45	2.63	263	1.39	1.39	1.39	0.92	2.7	2.7	6.89	66:0	1.14	0.99	2.9	1.9	1.9	
	ORF SEQ ID NO:	37235						32949					34977	35027			35190	35191	36026	36856	36857	36932	37776	. 37777	26774	27161	27162	29166	31735	27638	28067		32576	32577	
	SEQ ID NO:						18985	i I		1	20520		21812		21917	ı	22021	22021	22808	23606	23606	23676	24434	24434	4099	14462	14462	16527	18772	14905	15324	15458	19547	19647	
	Probe SEQ ID NO:	11280	11445	11445	881	1253	6210	7190	7286	7826	7825	9014	9124	9192	9238		9267	9267	10160	10926	10926	11004	11850	11850	1331	1/1B	1719	3//6	2881	2176	2813	2763	8847	6847	
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	Top Hit Descriptor	yo17g09.r1 Soares adult brain N2b5HB55Y Home saplens cDNA clone IMAGE:178240 57	Homo sapiens uncharacterized bone marrow protein BM039 (BM039), mRNA	AJ403124 3.4 (downregulated in larynx carcinoma) Homo sapiens cDNA clone i8	AJ403124 3.4 (downregulated in larynx carcinoma) Homo saplens cDNA clone I8	601673686F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3956517 5'	Human fumarase precursor (FH) mRNA, nuclear cene encoding mitochondrial protein, complete ods	Homo saplens (huc) mRNA, complete cds	801297955F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3628213 5'	Homo sapiens Ran GTPase activating protein 1 (RANGAP1), mRNA	601172658F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:3528134 5'	Homo seplens chramosome 21 segment HS21C002	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo saptens fatty-acid-Coenzyme A ligase, long-chain 4 (FACL4) mRNA	Homo sapiens attractin precursor (ATRN) gene, exon 16	Homo sepiens attractin precursor (ATRN) gene, exon 16	Homo sapiens protein tyrosine kinase 2 beta (PTK2B) mRNA	Homo sapiens PDZ domain-containing guanine nucleotide exchange factor I (LOC51735), mRNA	Homo sapiens phosphatidylinositol 3-kinase, class 2, alpha polypeptide (PIK3C2A) mRNA	Homo sapiens hypothetical protein FLJ10488 (FLJ10488), mRNA	Homo saplans hypothetical protein FLJ10488 (FLJ10488), mRNA	Homo sapiens SH3-domain GRB2-like 2 (SH3GL2), mRNA	Homo sapiens SH3-domain GRB2-like 2 (SH3GL2), mRNA	Homo saplens NKAT4b mRNA, complete cds	Homo sapiens NKAT4b mRNA, complete cds	H.sapiens arginase gene exon 3 (EC 3.5.3.1)	Homo sapiens AIM-1 protein (LOC51151), mRNA	Human cytochrame P450 (CYP2A13) gene, complete cds	M36b04x1 NCI_CGAP_Ut1 Homo sapiens cDNA done IMAGE:2281743 3' similar to SW:RL2B_HUMAN	P29316 60S RIBOSOMAL PROTEIN L23A;	PM0-BN0065-100300-001-c06 BN0065 Homo saplens cDNA	yc3105.r1 Soares fetal liver spleen 1NPLS Homo sapiens cDNA clone IMAGE:243585 5' similar to PIR:S64204 S54204 ribosomal protein L29 - human ;	
	Top Hit Database Source	EST HUMAN	N	EST HUMAN	EST_HUMAN	EST_HUMAN	IN	NT	EST_HUMAN		EST_HUMAN	۲N	ΙN	NT	N	LN L	N	LN	ΙN	LN	LN	IN	TN	LN	IN	NT	IN	IN		EST_HUMAN	EST_HUMAN	EST_HUMAN	
	Top Hit Acession No.	H46698.1	TN 9602268	AJ403124.1	AJ403124.1	BE900454.1	U59309.1	L26405.1	BE382519.1	11418177	BE294281.1	AL 163202.2	AF032897.1	4758331	AF218902.1	AF218902.1	4758975 NT	7706512 NT	4505798 NT	11431271 NT	11431271 NT	11428813 NT	11428813 NT	L76666.1	L76666.1	X12664.1	T705868 NT	U22028.1		AI862007.1	AW998611.1	N49818.1	
	Most Similar (Top) Hit BLAST E Value		3.0E-98	3.0E-98		3.0E-98	3.0E-98		3.0E-98	3.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98	2.0E-98		2.0E-98	2.0E-98	2.0E-98	_		1.0E-98 /	1.0E-98	
	Expression Signal	2.73	0.48	1.42	1.42	96.0	4.11	1.58	1.47	3.56	2.86	1.53	0.8	3.21	1.34	1.34	1.39	4.03	1.15	1.07	1.07	4.94	4.94	0.58	0.58	1.48	1.37	1.42		16.4	2.12	11.24	
	ORF SEQ ID NO:	34485	35030	35639	35640	36233	36799				27531							30757	32329			34342		34421		35290		37103		25829	25871	27237	
	Exan SEQ ID NO:	21341	21866	22434	L.	23017	23552	L	25262	25013									19322			21198						23822		_ [13228	14529	
	Probe SEQ ID NO:	8649	9197	9783	9783	10371	10872	11863	12668	12751	2071	2231	4264	4306	4778	4778	5131	5292	6557	7523	7523	8506	8506	8591	8591	9437	10313	11155		396	442	1789	

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Top Hit Descriptor	zp98c09.r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:628240 6' similar to TR:G806662 G806562 NEBULIN.;	601284986F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3606692 5	601284986F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3606692 5	Homo sapiens beta-tubulin mRNA, complete cds	Homo sapiens beta-tubulin mRNA, complete cds	QV-BT073-191298-012 BT073 Homo sapiens cDNA	QV-BT073-191298-012 BT073 Homo saplens cDNA	EST380711 MAGE resequences, MAGJ Homo saplens cDNA	tm89h07.x1 NCI_CGAP_Briz5 Homo sapiens cDNA clone IMAGE;2163421 3' similar to SW:bID_HUMAN P55957 BH3 INTERACTING DOMAIN DEATH AGONIST;	tm69h07.x1 NCI_CGAP_Bm25 Homo saplens cDNA done IMAGE:2163421 3' olmilar to SW:BID_HUMAN P55957 BH3 INTERACTING DOMAIN DEATH AGONIST;	zn90d02.r1 Stratagene lung carcinoma 937218 Homo sapiens cDNA clone IMAGE:565443 6' similar to TR:0662994 G682994 GPLANCHORED PROTEIN P137.;	Human endogenous retrovirus, complete genome	Homo sapiens oscillin (hLn) gene, exon 5	Hamo sapiens NK-receptor (KIR-G2) gene, linker region exon	Human G2 protein mRNA, partial cds	EST388473 MAGE resequences, MAGN Homo saplens cDNA	Homo sapiens CD34 antigen (CD34) mRNA	Homo sapiens GAP-like protein (LOC51308), mRNA	Homo sapiens polycystic kidney disease (PKD1) gene, exons 27-30	Homo sapiens polycystic kidney disease (PKD1) gene, exons 27-30	H.sapiens mRNA for estrogen receptor	Homo sapiens ankyrin-like with transmembrane domains 1 (ANKTM1), mRNA	Homo sapiens NDST4 mRNA for N-deacetylase/N-sulfotransferace 4, complete cds	Homo sapiens Iodestar protein mRNA, complete cds	Homo sapiens lodestar protein mRNA, complete cds		Homo sapiens inositol 1,4,5-triphosphate receptor, type 1 (ITPR1), mRNA			Hamo saplens UDP-glucose;glycoprotein glucosylfransferase 1 (HUGT1), mRNA	· ************************************
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝΤ	LN T	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	J	TN	LN		EST_HUMAN	TN	NT	NT	TN	NT	TN	NT	NT	NT	NT	LN L	TN	LN L	۲	
Top Hit Acession No.	\A195854.1			4F141349.1	41349.1	5004.1	AI905004.1	4W968635.1	4,1479829.1	4 479829.1	34604.1	9635487 NT	7.0E-99 AF035808.1	7.0E-99 AF001886.1	6.0E-99 U10991.1	4W976364.1	4502660 NT	7706136 NT	143610.1	43610.1	X99101.1	6601589 NT	AB036429.1	6.0E-99 AF080255.1	6.0E-99 AF080255.1	11431994 NT	11431994 NT	11526299 NT	9910279 NT	9910279 NT	
Most Similar (Top) Hit BLAST E Value	1.0E-98 AA1	1.0E-98	1.0E-98 BE3	1.0E-98 AF1	1.0E-98 AF1	9.0E-99 AI90	9.0E-99 AI90	9.0E-99 AW	9.0E-99 A147			8.0E-99	7.0E-99				6.0E-99	6.0E-99	6.0E-99 L43610.1	6-30.8					١					6.0E-89	 -
Expression Signal	3.4	1.1	1.1	2.7	2.7	0.88	0.88	4.21	2.75	2.75			10.03	2.99	1.89		1.21	0.72				0.53									
ORF SEQ ID NO:	30668	31178	31179	34728	34727	31441	31442	31691	37009	l		1			25886	١.		L									L			36432	
Exen SEQ ID NO:	18040	1	L	i_	21587	18520	1	Ì	23736	1	1	L	L	ı	١.	L	L	1	L	1		L	L	L	L	L	┸	1		L]
Probe SEQ ID NO:	5234	5482	5482	8898	8836	5728	5728	5949	11088	11086	11390	8627	5743	11610	469	3868	4699	6503	6578	6578	8003	8022	8663	8782	8762	8824	8	10820	11433	11433	

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xp09e06.x1 NCI_CGAP_HN9 Homo sepiens cDNA clone IMAGE:2739874 3' similer to gb:M31212 MYOSIN LIGHT CHAIN ALKALI, NON-MUSCLE ISOFORM (HUMAN); Homo sapiens short chain L-3-hydroxyacyl-CoA dehydrogenase precursor (HADHSC) gene, nuclear gene hd02h02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2908371 3' similar to TR:0027 zb46d06.r1 Soares_fetal_lung_NbHL19W Homo saplens cDNA clone IMAGE:306635 5' similar to Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA Homo sapiens FK506-binding protein 6 (36kD) (FKBP6) mRNA, and translated products Homo sapiens FK506-binding protein 6 (36kD) (FKBP6) mRNA, and translated products y81b09.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:145625 5 Homo saplens truncated Niemarin-Pick C3 protein (NPC3) mRNA, complete cds Homo sapiens truncated Niemann-Pick C3 protein (NPC3) mRNA, complete cds Homo saplens T cell receptor beta locus, TCRBV7S3A2 to TCRBV12S2 region Homo sapiens potassium channel, subfamily K, member 10 (KCNK10), mRNA Homo sepiens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds 601513157F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3914391 Human interferon-alpha receptor (HuiFN-alpha-Rec) mRNA, complete cds Homo sapiens intersectin long isoform (ITSN) mRNA, complete cds Homo sapiens cell recognition molecule Caspr2 (KIAA0868), mRNA gb:M15182 BETA-GLUCURONIDASE PRECURSOR (HUMAN); Human E2A/HLA fusion protein (E2A/HLF) mRNA, complete cds Homo saplens fatty acid amide hydrolase (FAAH) gene, exon 14 Homo sapiens fatty acid amide hydrolase (FAAH) gene, exon 14 Top Hit Descriptor Homo sapiens huntingtin interacting protein 1 (HIP1), mRNA Homo sapiens huntingtin interacting protein 1 (HIP1), mRNA Homo sapiens glycine receptor, alpha 2 (GLRA2), mRNA Homo sapiens glycine receptor, alpha 2 (GLRA2), mRNA Homo sepiens myosin X (MYO10) mRNA, complete cds Homo sapiens ALEX1 protein (LOC51309), mRNA Human Ku (p70/p80) subunit mRNA, complete cds Human Ku (p70/p80) subunit mRNA, complete cds 002711 PRO-POL-DUTPASE POLYPROTEIN; encoding mitochandrial protein, complete cds H.sapiens IMPA gene, exon 8 H.sapiens E6-AP gene exon HUMAN HUMAN HUMAN EST_HUMAN Top Hit Database Source EST EST_ RST EST 집 È 눋 눋 10863960 NT 11526150 NT 눋 눋 닐호 7427514 NT 4503730 NT z <u>'E</u> <u>5</u> 11421007 NT 11419721 NT 7427514 NT 11421007 Top Hit Acession 1.0E-99 AW340174.1 2.0E-99 AW274792.1 2.0E-99 AF247457.2 5.0E-99 BE890177.1 2.0E-99 AF095703.1 2.0E-99 108 1.0E-99 AF114487.1 1.0E-99 AF192523.1 .0E-99 AF 192523.1 I.0E-99 AF098018.1 ğ 1.0E-99 AF098018.1 2.0E-99 WZ3507.1 5.0E-99 Y11365.1 5.0E-99 AF009660 3.0E-99 M95586.1 2.0E-99 M30938. 2.0E-99 0E-99 1.0E-99 1.0E-99 1.0E-99 1.0E-99 1.0E-99 66-30. Aost Similar (Top) Hit BLAST E 1.56 4 8 8 8 8 9.55 6.88 8. 2.64 88. 1.02 2.82 1.71 2.01 2.64 4. 0.68 Expression Signal 27406 29874 37029 37030 28666 33348 25809 27367 27368 28489 35270 ORF SEQ ÖNQ 17261 24674 13968 22098 23754 14693 17241 13166 14295 14295 14657 15848 19622 SEQ ID 14657 17086 23754 20243 18453 2091 ö <u>8</u> 4526 1549 1920 8220 1217 3253 4506 8608 11049 1920 4347 6606 9420 11084 370 1549 4347 SEO ID 1957 8 6707 757

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Top Hit Descriptor	Homo saplens heat shock transcription factor 2 binding protein (HSF2BP), mRNA	Homo sapiens mRNA for KIAA1005 protein, partial cds	Homo saplens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, complete cds	Homo sapiens chromosome 21 segment HS21C047	Homo saplens chromosome 21 segment HS21C047	Homo sapiens Testis-specific XK-related protein on Y (XKRY), mRNA	Homo sapiens Testis-specific XK-related protein on Y (XKRY), mRNA	xv78b11.x1 NCI_CGAP_Bm53 Homo saplens cDNA clone IMAGE:2824605 31	Homo sapiens chromosome 21 segment HS21C006	Homo sapiens chromosome 21 segment HS21C049	EST02975 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCR32	Homo sapiens X-linked anhidrolitc ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	G.garilla DNA for ZNF80 gene homolog	RC3-HT0625-040500-022-b09 HT0625 Homo sapiens cDNA	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA	Homo sapiens DKFZP686M0122 protein (DKFZP586M0122), mRNA	602072064F1 NCI_CGAP_Brn67 Homo saplens cDNA clone IMAGE:4215039 5'	UI-H-BI1-afk-c-07-0-UI.s1 NCI_CGAP_Sub3 Home sapiens cDNA clone IMAGE:2722164 3'	qf62f09x1 Soares_testis_NHT Homo saplens cDNA clone IMAGE:1754633 3' similar to SW:CYT_COTJA P81081 CYSTATIN ;	Homo saplens mRNA for KIAA1168 protein, partial cds	Rat mRNA for short type PB-cadherin, complete cds	H.sapiens mRNA for IFN-gamma (pKC-0)	Homo sapiens KIAA0957 protein (KIAA0957), mRNA	Homo saplens RGH2 gene, retrovirus-like element	Homo sapiens myotubularh-related protein 1a mRNA, partial cds	Homo sapiens follide stimulating hormone receptor (FSHR) mRNA	Homo saplens myelin transcription factor 1-like (MYT1-I) mRNA, complete cds	Homo saplens small optic lobes (Drosophila) homolog (SOLH) mRNA		601863164F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE: 4080999 5'.	the second
Top Hit Database Source	N	TN		뉟	NT	IN	ΙΝ		EST HUMAN	NT	NT	EST_HUMAN		N	NT	EST_HUMAN	NT	IN	EST_HUMAN	EST HUMAN	EST_HUMAN	Ę	NT	TN	LN	F	NT	NT	ΙN	LN	NT	EST_HUMAN	
Top Hit Acession No.	5901979 NT	1.0E-99 AB023222.1		1.0E-99 AF240786.1	NL163247.2	AL163247.2	11418230 NT	11418230 NT	W275237.1	NL163206.2	AL163249.2	05087.1		1.0E-100 AF003528.1	(89631.1	1.0E-100 BE180609.1	7661685 NT	7661685 NT	1.0E-100 BF530735.1	1W207555.1	1.0E-100 AI200857.1	1.0E-100 AB032994.1	J83349.1	(62468.1	11418976 NT	011078.1	05735	4503792 NT	1.0E-100 AF036943.1	5032104 NT	5032104 NT	1.0E-100 BF244218.1	
Most Similar (Top) Hit BLAST E Value	1.0E-99	1.0E-99 /		1.0E-99/	1.0E-100 AL	1.0E-100 AL	-1.0E-100	1.0E-100	1.0E-100 AW	1.0E-100 AL	1.0E-100 AL163249	1.0E-100		1.0E-100/	1.0E-100 X89631.1	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100 /	1.0E-100	1.0E-100 /	1.0E-100 D83349.1	1.0E-100	1.0E-100	1.0E-100 D11078.1	1.0E-100 AF	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	
Expression Signal	1.8	2.77		6.68	. 0.95	1.53	1.54	1.54	0.69	1.24	0.83	3.08		1.28	7.19	1.33	3.22	3.22	3.14	1.14	1.81	1.41	1.39	1.33	2.36	5.5	1.52	2.14	1.03	2.66	2.66	1.62	
ORF SEQ ID NO:	37088	37343			25443	25443	25526	25527	25550	25618	25763	25770				25915	26418	26419	26846		26976	27309		27894	28166		29558	29578	29785	30362	30363	30637	
Exon SEQ ID NO:	23808	24040		24526	12830	12830	12894	12894	12911	12978	13113	13136		13213	13263	13280	13758	١.	1	14285		1	ł	15159	l	15784	16927	L	l	1	l	1	
Probe SEQ ID NO:	11141	11350		11984	-	2	88	99	85	165	88	334		427	477	496	866	88	1415	1538	1543	1856	2238	2439	27.10	3018	4188	4211	4418	5031	5031	5207	

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	Top Hit Descriptor	xa82f01.x1 NOI_CGAP_CML1 Homo sapiens cDNA clone IMAGE:2573305 3' similar to gb:X12433 PROTEIN PHPS1-2 (HUMAN);	AU118182 HEMBA1 Homo sapiens cDNA clone HEMBA1003046 5'	Homo sapiens NF-E2-related factor 3 gene, complete cds	Human mRNA for plasma inter-alpha-trypsin inhibitor heavy chain H(3)	Homo sapiens ER to nucleus signalling 1 (ERN1) mRNA	Homo saplens ER to nucleus signalling 1 (ERN1) mRNA	Homo sapiens hect domain and RLD 2 (HERC2), mRNA	AU140214 PLACE2 Homo sapiens cDNA clone PLACE2000137 5'	AU136800 PLACE1 Hamo sapiens cDNA clane PLACE1005089 5'	J/38c08.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:129134 3'	Homo sapiens Rho GTPase activating protein 8 (ARHGAP8), transcript variant 4, mRNA	ae33b06.r1 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:897687 5' similar to TR:G487418 G487418 ACTIN FILAMENT-ASSOCIATED PROTEIN.;	ae33b06.r1 Gessler Wilms tumor Homo sapiens cDNA clone IMAGE:897587 5' similar to TR:G487418	1646/418 ACTIN FILAMEN I-ASSOCIATED PROTEIN;	AND STATE OF THE S	MKT-IN0046-060900-004-b05 TN0046 Homo saplens cDNA	Human mRNA for kidney epidermal growth factor (EGF) precursor	601647357F1 NIH_MGC_61 Homo sapiens cDNA clone IMAGE:3931310 5'	Homo saplens chromosome 21 segment HS21C003	AU116961 HEMBA1 Homo sapiens cDNA clane HEMBA1000343 5'	AU116951 HEMBA1 Homo sapiens cDNA clone HEMBA1000343 5'	Homo sapiens mRNA for KIAA1485 protein, partial cds	wr37g09.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2489920 3' similar to contains element	Micraz repended element;	FMM-SNUGGO-100300-001-006 BNUGGS Homo Sapiens CDNA	AU12/120 N 12KPZ Hamo sapiens cDNA clone N 12KP2001918 5	Homo saplens mRNA for KIAA1626 protein, partial cds	Homo sapiens mRNA for KIAA1628 protein, partial cds	hh83c11.y1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2969396 5	IMAGE:2969396 5'		602020554F1 NCI_CGAP_Brn67 Homo septens cDNA clone IMAGE:4156165 5'
	Top Hit Database Source	EST HUMAN	EST_HUMAN	F	Ę	Ę	FZ	攴	EST_HUMAN	EST_HUMAN	EST_HUMAN	17	EST_HUMAN	1487	EST HUMAN	N I I	EST HUMAN	LV.	EST HUMAN	LN⊤	EST_HUMAN	EST_HUMAN	NT	14441111	EST HOMAN	ES HUMAN	ESI HUMAN	L	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
	Top Hit Acession No.	075983.1	118182.1	1.0E-100 AF135116.1		4657568 NT	4557568 NT	5729867 NT	140214.1	136800.1	3887.1	7382479 NT	1.0E-100 AA496841.1	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	496841.1	1	_		-	63203.2		1.0E-100 AU116951.1			1	998011.1	1		46846.1	630487.1		1.0E-100 AV732101.1 E	47519.1
	Most Similar (Top) Hit BLAST E	1.0E-100 AW	1.0E-100 AU	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100 AU	1.0E-100 R1	1.0E-100	1.0E-100	707	1.0E-100 AA	ייייייייייייייייייייייייייייייייייייייי	1.0E-100	1.0E-100	1.0E-100	1.0E-100 AL1	1.0E-100	1.0E-100	1.0E~100	10.4	1.05-100	1.0E-100	1.0E-100 AU1	1.0E-100	1.0E-100	1.0E-100 AW	1.0E-100 AW	1.0E-100	1.0E-100 BF3
	Expression Signal	0.59	1.33	1.26	0.8	0.94	0.94	1.67	5.64	1.97	1.37	6.0	1.19	7,70	1.19	2 9	1.13	6.76	7.17	4.8	0.68	0.68	3.62	ç	2.78	1.82	<u>رة.</u>	2.11	2.11	1.68	1.68	0.49	1.47
	ORF SEQ ID NO:	30831	31323	31369	31461	31814	31815		32173	32224	32362	32622	32608	00000	32009	10070	32558	32565	34261		34746	34747	34986		202.00	33320	$\frac{1}{1}$	35316	35317	35598	35599	35749	36230
	Exan SEQ ID NO:	18220	18410	18455	18539	18850	18850	19113	١.	19224	19349	19587	19576	40676	1	H	-	-1	21123			21603		0000	800	40404	2	22136	22136	22394	22394	22554	23014
	Probe SEQ ID NO:	5421	5614	2660	5747	6071	6071	6343	6405	6457	6586	6670	6742	6770	678R	3 5	08/0	6/93	8430	8466	8912	8912	9132	2	2 28	3 8	282	9483	9483	9743	9743	9905	10368

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Zk89a03.rf Soares_pregnent_uterus_NbHPU Homo sapiens cDNA clone IMAGE:489964 67
Zk89a03.rf Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:489964 67
Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) 7488h03.x1 NCI_CGAP_Lu24 Home sapiens cDNA clone IMAGE: 3' similar to TR:Q21897 Q21897 COSMID R151. [2] TR:Q9UA08; Homo saplens phosphoribosy/glycinamide formyfransferase, phosphoribosy/glycinamide synthetase, Homo sapiens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA 602168474F1 NIH_MGC_83 Homo sepiens cDNA clone IMAGE:4297291 6' q999e09x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1843336 3' Homo sepiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA Homo sepiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene Homo sapiens 14q32 Jagged2 gene, complete ods; and unknown gene Homo sapiens SH3-domain binding protein 1 (SH3BP1), mRNA Homo sapiens transcobalamin II: macrocytic anemia (TCN2), mRNA Homo sepiens A kinase (PRKA) anchor protein 6 (AKAP6), mRNA Top Hit Descriptor Homo sapiens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mRNA Homo sapiens SEC14 (S. cerevisiae)-like 2 (SEC14L2), mRNA Homo sapiens mRNA for KIAA0448 protein, partial cds Homo seplens chromosome 21 segment HS21C047 QV2-PT0012-010300-070-d04 PT0012 Homo seplens cDNA 270300-008-h11 BN0070 Homo saplens cDNA Homo sapiens carboxypeptidase A1 (pancreatic) (CPA1) mRNA Homo saplens golgin-like protein (GLP) gene, complete cds RC3-ST0281-160600-016-h09 ST0281 Homo saplens cDNA Homo sapiens KIAA0569 gene product (KIAA0569), mRNA Homo sapiens ventral anterior homoobox 2 (VAX2), mRNA Homo saplens KIAA0569 gene product (KIAA0569), mRNA Homo sapiens ventral anterior homeobox 2 (VAX2), mRNA phosphoribosylaminoimidazole synthetase (GART) mRNA Homo sapiens of cardiac alpha-myosin heavy chain gene Human endogenous retrovirus HERV-K, pol gene Single Exon Probes Expressed in Brain H.sapiens EWS gene, exon 5 genes, complete cds EST_HUMAN Top Hit Database Source **EST HUMAN** EST_HUMAN EST HUMAN EST_HUMAN HUMAN ż 7110714 NT 7110714 NT 11417974 NT Ξ Ł 7110734 NT 4503914 NT 7662183 NT 7662183 NT 7657454 NT 5729892 NT Top Hit Acession 11545732 5921460 5921460 4502996 1.0E-100 AW875464.1 1.0E-100 AF266285.1 1.0E-100 AL163247.2 1.0E-100 AF111170.3 .0E-100 AA115605.1 1.0E-100 AF240786.1 1.0E-100 BF 446549.1 AB007915.2 1.0E-100 AA115605. BF681218.1 1.0E-101 X72993.1 1.0E-100 Most Similar (Top) Hit BLAST E Value 1.0E-100 1.0E-100 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 1.0E-101 4.52 6.27 4.52 2.11 1.59 1.48 1.57 6.67 1.75 5.88 1.57 3.67 Expression 1.62 5.88 1.22 1.52 1.52 1.52 1.63 1.5 1.71 Signal ORF SEQ 37319 37320 25443 37668 37806 30968 25539 25540 26087 26111 ÖΝΩ 26112 26273 26396 26449 26185 27012 27183 SEQ ID NO: 24017 12830 24340 24471 13463 12902 25044 13447 13729 13526 13671 14324 14324 14482 14482 14673 15332 Probe SEQ ID 10658 11326 1356 10458 11633 11749 11907 11681 12031 12792 688 688 754 1030 ġ 671 833 4746 1938 2349 1577

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Top Hit Descriptor	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens genomic downstream Rhesus box	Hamo sapiens gamma-glutamyltransferase 1 (GGT1) mRNA	601458531F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3862086 5'	EST377629 MAGE resequences, MAGI Homo sapiens cDNA	Homo saplens RiBilR gene (partial), exon 12	Homo sapiens RiBliR gene (partial), exon 12	Homo sapiens ASH2L gene, complete cds, similar to Drosophila ash2 gene	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	EST377212 MAGE resequences, MAGI Homo sepiens cDNA	Homo sapiens cytoplasmic linker 2 (CYLN2), mRNA	Hamo sapiens cytoplasmic linker 2 (CYLN2), mRNA	Homo sapiens carbonic anhydrase VII (CA7), mRNA	Homo sapiens hypothetical protein FLJ22087 (FLJ22087), mRNA	Homo sapiens Kruppel-type zinc finger protein (PEG3) mRNA, alternative splice form 4, partial cds	Homo sapiens Kruppel-type zinc finger protein (PEG3) mRNA, alternative splice form 4, partial cds	ws5ff2.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2533487 31	601109217F1 NIH_MGC_16 Hamo sapiens cDNA clone IMAGE:3349901 5'	RC1-BT0313-220700-018-f12 BT0313 Homo sapiens cDNA	601121621F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3345869 5'	601121621F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3345869 5	601764686F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3996837 5	hh74g10.71 NG_CGAP_GU1 Homo sapiens cDNA done IMAGE:2988578 5' similar to gb:J03143 INTERFERON-GAMMA RECEPTOR ALPHA CHAIN PRECURSOR (HUMAN);	hh74g10 y1 NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:288878 5' similar to gb:J03143 INTERFERON-GAMMA RECEPTOR ALPHA CHAIN PRECURSOR (HUMAN);	zk29g08.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:471998 5' stmflar to	PIR:S54640 S54640 YD9335.03c protein - yeast;	Homo saplens mRNA for KIAA1351 protein, partial cds	Homo saplens mRNA for KIAA1351 protein, partial cds	Human mRNA for pancreatic gamma-glutamyltransferase	Human mRNA for pancreatic gamma-glutamyltransferase
Top Hit Database Source	N	ΙN	NT	NT	EST_HUMAN	EST_HUMAN	TN	ŢN	LN	LN	LN.	EST_HUMAN	LN	TN	NT	ΙΝ	LN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN	NT	L	NT	NT
Top Hit Acession No.	1.0E-101 AJ237744.1	1.0E-101 AJ237744.1	1.0E-101 AJ252312.1	4885270 NT	1.0E-101 BF035327.1	1			1.0E-101 AB022785.1	5921460 NT	5921460 NT	4W965139.1	1.0E-101 7427512 NT	7427512 NT	11430734 NT	11545780 NT	1.0E-101 AF208970.1	1.0E-101 AF208970.1	1.0E-101 AW008475.1				1.0E-101 BE275821.1	1.0E-101 BF029174.1	1.0E-101 AW630070.1	AW630070.1				AB037772.1		X60069.1
Most Similar (Top) Hit BLAST E Value	1.0E-101	1.05-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101		1.0E-101	1.0E-101	1.0E-101		1.0E-101)
Expression Signal	1.09	1.09	13.73	1.98	2.27	1.93	1.49	1.49	3.69	1.16	1.16	1.22	3.68	3.68	1.27	1.01	5.57	29.57	7.48	1.79	7.43	0.84	0.84	2.88	0.67	0.67		1.08	0.83	0.83	17.13	17.13
ORF SEQ ID NO:		28193		28613	,	28790			29245	30304	30305			31652	32372				33136					33772	34050	34051	-		35080			33554
Exon SEQ ID NO:	15452	15452	15721	15961	15997	16134	15452	15452	16607	17697	17697	18041	18698	18698	19358	19859	19905	19905	20058	20147	20289	20508	20508	20649	20915	20915		21599	21908	21908	20432	20432
Probs SEQ ID NO:	2747	2747	2925	3198	3235	3375	3395	3395	3857	4974	4974	5235	5913	5913	6595	7173	7220	7220	7376	7474	7623	7813	7813	7954	8221	8221		8908	9229	9229	9362	9362

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Top Hit Descriptor	Homo sapiens gamma-glutamyltransferase 1 (GGT1), transcript variant 3, mRNA	60147280811 NIH_MGC_68 Homo saptens cDNA clone IMAGE:3876953 3	601472808T1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3875953 3	Homo sapiens potassium channel, subfamily K, member 10 (KCNK10), mRNA	Homo sapiens Janus kinase 2 (a protein tyrosine kinase) (JAK2), mRNA	601680825F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950887 5'	601680825F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:3950887 5	branched-chain alpha-keto acid dehydrogenase complex E1 alpha subunit [human, Genomic, 195 nt, segment 8 of 9]	Homo sapiens mRNA for KIAA0819 protein, partial cds	tm58c01.x1 NCI_CGAP_Bm25 Homo sapiens cDNA cione IMAGE:2182304 3' similar to gb:M13361 HEPARIN-BINDING GROWTH FACTOR PRECURSOR 1 (HUMAN);	tm68c01.x1 NCI_CGAP_Brn25 Homo saplens cDNA done IMAGE.2162304.3° similar to gb:M13391 HEPARIN-BINDING GROWTH FACTOR PRECURSOR 1 (HUMAN);	RC-BT163-290499-085 BT163 Homo squiens cDNA	RC-BT163-290499-085 BT163 Homo sapiens cDNA	QV1-DT0068-240200-085-a01 DT0068 Homo sapiens cDNA	Homo sapiens phosphatidylinositol 4-kinase 230 (pl4K230) mRNA, complete cds	Homo saplens chromosome 21 segment HS210103	Homo sapiens down-regulated in adenoma (DRA) mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo sapiens southe carrier taminy ((adminated glucose utarspones), manipula et olocose principula (manipula et olocose), misera	Homo sapiens solute carrier tamily 2 (racilitated glucose transporter), member 9 (35,02,04,0), minor	Homo saplens reelin (KELN) mKNA	601289982F1 NIH_MGC_21 Home sapiens conveniented as a second of second s	am60c10.x1 Johnston frontal cortex Homo sapiens cDNA cione iMAGE:1339934.3 similar to SW:GG95_HUMAN Q08379 GOLGIN-95.;	am@0c10.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539954 3' similar to	SW. GGGS TICMAN GGS/8 GOLOIN-50.	Training sapitatis nitrator to game product (nitrator to f), minator	AU141000 PLACE4 nome sapiens contact district solutions of the place o	AU141005 PLACE4 nomo sapiens conva done reaccessos s	Homo sapiens chromosome 21 segment no.21 Court	60TTU/64SFT NIT MIC TO FOUR Equipments of the control of the contr
Top Hit Database Source	I. I	EST_HUMAN	EST_HUMAN	ΤN	TN	EST_HUMAN	EST_HUMAN	Þ	Į,	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	NT	NT	ΓN	LN.	NT	N	EST_HUMAN	EST_HUMAN		EST_HUMAN	- 1	EST HUMAN	EST_HUMAN	LN I	EST HUMAN
Top Hit Acession No.	9845492 NT	BE619667.1	BE619667.1	10863960 NT	11429127 NT	BE973648.1	BE973648.1	S3837.1	AB020626.1	AI590078.1	A1590078.1	A1908168.1	Al908168.1	AW939051.1	1.0E-102 AF012872.1	1.0E-102 AL 163303.2	4557534 NT	M10976.1	11437146 NT	11437146 NT	4826977 NT	2 BE408447.1	1.0E-102 A1124669.1		1.0E-102 AI124669.1	7661979 NI	1.0E-102 AU141005.1	1.0E-102 AU141005.1	1.0E-102 AL163207.2	BE251310.1
Most Similar (Top) Hit BLAST E Value	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1=	1.0E-101		1.0E-101		$\overline{}$. 1			1.0E-102			1.0E-102 M10976.	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102		1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102	1.0E-102
Expression Signal	17.01	6,24	6.24	0.72	1.49	9.0 19.0	26:0	108	2.11	2.06	2.06	1.31	1.31	13.68	2	4.35	1.59	2.8	1.67		66'0	119.7	1.88				4.76			2.17
ORF SEQ ID NO:	35123		1										L		26485	25767			26660		26678		27769			28472				29748
Exen SEQ ID NO:	21950	1	L	1	22955	•	L		_	ł	ل	L	L	L		13133	1_	L	13994	13894	14010		15032	.1		15827	15895	15895		17115
Probe SEG ID NO:	9375	9657	9857	9794	10308	10447	10447	0040	11053	11308	11308	11783	11763	12461	38	332	758	1095	1245	1245	1261	1398	7307		2307	3061	3130	3130	4207	4378

Page 427 of 536 Table 4 Single Exon Probes Expressed in Brain

			_	- 1	_	_	_	-		П	_	П	ГТ			П			_		Ĭ	П	٦	٦	٦		T	T				
	Top Hit Descriptor Top Hit Descriptor	Homo sapiens protein phosphatase- I regulated y success. Homo sapiens protein phosphatase- I regulated y success.	Homo sapiens HSC34 IIINN 19 TO TAIL A	Homo sapidas historie deadeaptate i (1970), Paris Pina Pina Pina Pina Pina Pina Pina Pina	Homo sapiens histone deacetytase / (http://www.new.	Homo sepiens hect domain and RLD 2 (HERCZ), mixed Activities 31 similar to TR:Q13137	ar82f09.x1 Barstead colon HPLRB7 Homo sapiens culva ciolie ivinos	Q13137 NDP52. ; 801561505F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3831241 5"	601277215F1 NIH_MGC_20 Homo sapiens culva dione invacations	Homo saplens mRNA for Centaurin-alpha2 protein	AV710738 Cu Homo sapiens cUNA cigne Curvincus Cona	QV3-NT0025-210600-236-h08 NT0025 Home saptens COV3-NT0025-210600-236-h08 NT0025 Home saptens COV3-NT0025-210600-236-h08 NT0025 Home saptens COV3-NT0025-3003145 51	601501107F1 NIH_MGC_70 Homo saplens cLINA cione ilwayccoc.	AV694817 GKC Homo sapiens cDNA clone GNOEEF 11 3	AV694817 GKC Homo saplens cDNA clone GKCEEE 1.3	Homo sapiens mRNA for KIAA0454 protein, partial cus	601283770F1 NIH_MGC_44 Homo saplens cDINA close IMAGE:3605536 5	601283770F1 NIH MGC 44 Homo sapiens CONA clone IMAGE 2397971 3' similar to contains MER4.tf	Wig3b06.x1 NCI_CGAP_Kld12 Homo sapiens con con-	MER4 MER4 repatitive element : AV755842 BM Homo sapiens cDNA clone BMFAUD08 5'	vd13d07.r1 Soares fetal liver spleen 1NFLS Homo sapiration control of the liver spleen 1NFLS control of the	1			Г	Г	Г	$\neg \Gamma$	Homo sapiens UDP glycosyltransferase 2 family, polypeptuse 211 (1993473)	Г	1 1	
	Top Hit Database Source	Z	Z		15	L		EST_HUMAN	EST HIMAN	L	EST HIMAN	EST HI IMAN	EST HUMAN	EST HIMAN	EST HIMAN	LN	FST HUMAN	EST HUMAN		EST HUMAN	EST HIMAN	EST HUMAN	EST HUMAN	Z	EST HUMAN	EST HUMAN		EST_HUMAN	TNIC	EST HIMAN	EST_HUMAN	
5	op Hit Acession No.	F067133.1		5398	TVI ROSSONT	TIM SACCOLAR	11433040			A 1038004 4	740700 4	AV/10/30.1	BE/63051.1	E910555.1	AV694817.1	AV094017.1	AB00/923.1	BE388063.1		AI762859.1	1.0E-102 AV /55642.1	1 /0393.1	A11424820 1	1.0E-102/AD12-025	A1005037 1	1.0E-102/Alabour 1	Alexandr.	1.0E-102 AA970786.1 ES		10000	1.0E-102 AA8686/5.1	
}	Most Similar (Top) Hit BLAST E Value	1 0E-102 AF	112		1.DE-102	1.0E-102	1.0E-102			1.0E-102 B	1.0E-102/	1.0E-102	1.0E-102	1.0E-102	1.0E-102/			1.05-102	1.0E-10Z	1.0E-102	1.0E-102	1.0E-102 170393.1	100-102	1					1	1.0E-10	1.0E-10	
	Expression (Signal	- 6,	ò	9.1/	2.84	2.84	0.81	2.93	0.75	1.04	8.23	2.48	3.91	1.32	2.21	2.21	1.19	0.63	0.63	0.52	0.81	2.28	2.20	3.79			3.67	1.24			1.55	
	ORF SEQ E	-	30753		31414	31415	31420	31954	32785	32821	33014	33288		34032	34218	34219	34335		34668	35049	35017			35151			3 36229	36295	36937	36938		37 37287
	SEQ ID	1	18092	18458	18492	18492	18498	18976	19728	19756	19939	20195	1	١.	1	L	L	L	21521	<u> </u>	L	<u>L</u>	21900	<u> </u>	1 22929	L	Ш	23074	1	1_		30 23987
	Probe SEQ ID		5287	5683	2698	5608	5704	6200	7038	7065	7266	7524	8122	8201	8392	8392	8501	8828	8829	9450	918	9221	9221	9311	10281	10367	10367	1045	11008	41008	11290	11380

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1/57275																					_			P	C'	T/ l	US	01	<u>/0</u> 0	<u>66</u>	7	 .
Top Hit Descriptor	Human chromosome 16 creatine transporter (SLC6A8) and (CDM) paralogous genes, complete cde	Homo sapiens chromosome 21 segment HSZ10080	xk07c12x1 NCI_CGAP_CG20 Home septens course invoce account of the course	601500405F1 NIH_MGC_70 Homo sapiens cUNA cione invance: 3802400 3	601500405F1 NIH MGC 70 Homo sapiens cunna cinna livia de la constanta de la co	Home sapiens mRNA for KIAA0235 protein, partial cus	Homo sepiens nucleolar protein (NNC/D repeat) (NOF-30) IIINNA TAIN 6 - TAIN	Homo sapiens mixiva for programicy associated process – Community of the C	6014603866T INIT MICCOS TOTIO SEPTETE CONTINUE TO THE COMPIETE COS	Home sapiens priospriating in the responsibility of the responsibi	norm sequence sure control of sequence of the	Homo sapiens bone morphogeneuc protein 9 (ostoogenin protein 2) (1977) 3 (1978)	Homo sapiens bone multiplication process of contract of the co	AU134891 PLACET nome september out a construction of the complete cds	Home septemb promyter barkering and might promise an Indiana Charlette (179429 6)	COZZA 1002T I NO. COZA PINEZ Homo saniens CDNA clone IMAGE:4179429 5	SOCIATION OF THE MODE BENEFIT OF THE SAME SAME SOUND S	10(12) 13 13 14 14 14 14 14 15 15 15 15 15 15 15 15 15 15 15 15 15	UI-H-6W 0-girn-II-0-01/8-1WO COM COMMON COMM	Total organism of the state of	Macaca Intrigues Sychologics (#037240) Home sapiens cDNA clone IMAGE:840407 3' similar to contains	et louizat orangona marginata.	Homo saplens neuropilin 1 (NRP1), mRNA	seq340 b4HB3MA-Coff09+10-Bio Homo sapiens oDIVA clone b4nB3MA-Coff09-19-20-20	602186023-1 Nid-Mid-49 handra september 5 control of the control o	Homo sapiens septin 2 (SEP2) mKNA, parial cas	Homo sapiens KIAA0440 protein (KIAA0440), mKIVA	Homo sapiens KlAA0440 protein (KlAA0440), mKNA	EST36636 MAGE resequences, MAGC from seprens conv	EST386636 MAGE resequences, was continue appears of the 1301452 3'	ag/26e03.51 Soares testis Inn I multi bequetis conviction source to the confidence alimina recentor which 2 subunit (GLRA2) gene, exon 4	
Top Hit Database Source	TN	NT	EST HUMAN	EST_HUMAN	EST_HUMAN	N _T	Ļ.Z	L.	EST HOMAN	5	LN.	LN.	LZ.	EST_HUMAN	L _N	ESI HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N	Z	EST_HUMAN	NT	EST HUMAN	EST_HUMAN	NT	N⊤	SINT	EST_HUMAN	EST_HUMAN	EST HUMAN	Z
Top Hit Acession No.	141302.1	4L163280.2	4W300862.1	1.0E-103 BE908158.1	1.0E-103 BE908158.1	078.	5453793 NT	1.0E-103 AJZ78348.1	BE877541.1	1287	7657592 NT	. 4502428 NT		AU134991.1	AF060568.1	1.0E-103 BF528379.1	1.0E-103 BF528379.1	1.0E-103 BE744722.1	AW 298245.1	1.0E-103 AB040892.1	1.0E-103 AF023861.1	1.0E-103 AA485663.1	11430876 NT	1.0E-103 T23683.1	1,0E-103 BF569527.1	1.0E-103 AF179995.1	11435053 NT	11435053 NT	1.0E-103 AW954566.1	1.0E-103 AW954566.1	1.0E-103 AA781442.1	AF053490.1
Most Similar (Top) Hit BLAST E Value	1.0E-102 U41	1.0E-102 AL1	1.0E-102 AW	1.0E-103	1.0E-103	1.0E-103 D87	1.0E-103	1.0E-103			1.0E-103	1.0E-103	1.0E-103											Ľ		L			1			
Expression Signal	3.67	3.52	4.69	1.19	1.19	10.5	1.45	0.79	10,23		1.75	1.44	1.44	1.15				2.68			8.55	123			96'0					0.75		0.94
ORF SEQ ID NO:	37631		31023					26389	26643	27026	27354			27765	27909	28055	28058			28837		20168	l	L					L			32304
Exen SEQ ID NO:	24305	1	24820	12895	12895	L	L	L	13971	14337	14844	L	14704	15028	L	15316	15316	16830	16133	16189	16490	16528	L	L	1_	L	1	1	1_		1.	18301
Probe SEQ ID NO:	11710	11911	12450	87	87	86	203	960	1221	1591	1907	1968	1968	2303	2452	2603	2603	3064	3374	3433	3737	74.6	3810	3986	5844	5852	6174	8174	838	6368	6498	6535

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. Top Hit Descriptor	tm58b05.x1 NCI_CGAP_Bm26 Homo sapiens cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS. ;	tm58b05.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;	Homo sapiens dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS230, DXS239, DXS268, DXS269, DXS270, DXS272 (DMD), transcript variant Dp427m, mRNA	Homo saplens dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS230, DXS239, DXS288, DXS269, DXS270, DXS272 (DMD), transcript varient Dp427m, mRNA	Homo sapiens ribosomal protein L3-like (RPL3L), mRNA	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	EST377849 MAGE resequences, MAGI Homo sapiens cDNA	601571537F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:3838545 5'	tm58b05.x1 NCI_CGAP_Brn25 Homo sepiens cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;	tm58b05.x1 NCI_CGAP_Bm25 Home sapiens cDNA clone IMAGE:2162289 3' similar to TR:Q13769 Q13769 ANONYMOUS.;	EST27193 Human Brain Homo saplens cDNA 5' end similar to None	AU140344 PLACE2 Homo sapiens cDNA clone PLACE2000374 5'	AU140344 PLACE2 Homo sapiens cDNA clone PLACE2000374 5'	7160e.03.x1 Soares_NSF_F9_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3525964.3' similar to SW:PTNF_HUMAN Q16825 PROTEIN-TYROSINE PHOSPHATASE D1;	Homo sepiens triple functional domain (PTPRF Interacting) (TRIO), mRNA	Homo sapiens triple functional domain (PTPRF Interacting) (TRIO), mRNA	nd13c02.s1 NCI_CGAP_Ov1 Homo sapiens cDNA clone IMAGE:800162 3' similar to gb:L02426 26S PROTEASE SUBUNIT 4 (HUMAN);	ae84d12.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:970871 3' similar to gb:X03747 cds1 SODIUM/POTASSIUM-TRANSPORTING ATPASE BETA-1 (HUMAN);	QV2-NN0045-230800-322-b03 NN0045 Homo sapiens cDNA	QV2-NN0045-230800-322-b03 NN0045 Homo sepiens cDNA	H.sapiens mRNA for latent transforming growth factor-beta binding protein (LTBP-2)	EST375749 MAGE resequences, MAGH Homo sapiens cDNA
Top Hit Database Source	EST_HUMAN	EST_HUMAN	NT	NT	LN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	NT	LN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN
Top Hit Acession No.	0E-103 AI590071.1	1.0E-103 AI590071.1	5032282 NT	5032282 NT	11431100 NT	1.0E-103 AJ289880.1	1.0E-103 AW965776.1	1.0E-103 BE748158.1	1.0E-103 AI590071.1	1.0E-103 AI590071.1	T31080.1	4U140344.1	1.0E-103 AU140344.1	1.0E-103 BF109244.1	6005921 NT	6005921 NT	1.0E-103 AA581086.1	1.0E-103 AA774980.1	1.0E-103 BE935842.1	1.0E-103 BE935842.1	1.0E-103 Z37976.1	\W963676.1
Most Similar (Top) Hit BLAST E Value	1.0E~103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103	1.0E-103
Expression Signal	1.48	1.48	1.53	1.53	1.27	0.99	1.83	6.93	4.21	4.21	0.77	2.22	2.22	1.1	2.86	2.86	1.16	5.04	0.56	0.58	1.44	1.89
ORF SEQ ID NO:	32396	32397	30579	30580		32716		32990	33448	33447	34019	34354	34355		34845	34846	34891	34941			35811	35859
Exen SEQ ID NO:	19380	19380	17916	17916	ı	ĺ	19815	19918	20335	20335	1			21298	21695	21695	21736	21777	1	21832	22606	22647
Probe SEQ ID NO:	8618	6618	6747	6747	6872	6935	7127	7233	7871	7671	8187	8519	8519	8604	9005	9005	9046	9088	9162	9462	9928	6666

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Table 4 Single Exon Probes Expressed in Brain	Top Hit Descriptor	au51g04.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2618328 6' similar to	ol02d08,56 NCL_CGAP_Lu5 Homo sepiens cDNA clone IMAGE:1522283 5' sImilar to TR:062084 OB2084	FINOSPHOLIPASE C NEIGHBORING;	Home septems AXI	Home seriens NOB4 - 44:- Alons	Home series NOD1 protein (NOD1) gene, exons 1, 2, and 3	Home septems mON forcem (NOUT) gene, exons 1, 2, and 3	Home series mb/ly for partal OCT/plexin-A2 protein	Altagase bit Activities between OCI/plexin-A2 protein	Homo saplens polyment tuta.	Te88a10.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Home saplens cDNA clone IMAGE-32878410.9* climited.	contains MER29.t3 MER29 repetitive element;	Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA	Homo sepiens gene for AF-6, complete cds	DKFZp564H1072_r1 564 (synonym: hfbr2) Homo sapiens cDNA clone DKFZp564H1072 5	UNTZD564H1072_r1 564 (synonym: hfbr2) Homo saplens cDNA clone DKFZp564H1072 6	MAGE 887828 2' ole	gb:214116_ma1 CD59 GLYCOPROTEIN PRECURSOR (HUMAN);	601577460F1 NIH_MGC_9 Home saplens cDNA clone IMAGE:3926438 5	RC1-C10249-110900-214-f12 CT0249 Homo sapiens cDNA	NO (-0.10249-110900-214-f12 CT0249 Homo sapiens cDNA	Home september ANT Z (acum-telated protein 2, yeast) homolog (ACTR2), mRNA	Home Septens KlyA0440 protein (KIAA0440), mRNA		tplete cds		Homo seniens mRNA 64 KIA 8 1376			
igle Exon Pro	Top Hit Database Source	EST_HUMAN	TOUR LOUD	-1.	Į	Ļ	L	LN L	LN	EST HIMAN			ESI HOMAN			EST_HUMAN	ES L'HOMAN		EST HUMAN	ESI HUMAN	EST HOMAN	NAMOL 101					T HUMAN				- LN
Sir	Top Hit Acession No.	A1878956.1	1792759 1	11424061 NT	11424061 NT	1467	1.0E-103 AF149773.1	37831.2	87831.2	Γ			44500004	1020201	097540.0	03/549.3	4503426	02470	1	T	34224.1	5031570	7882125 NT	7682125 NT	671.1	151.1		Γ		032998.1 NT	
	Most Similar (Top) Hit BLAST E Value	1.0E-103 A	1.0E-103 A	1.0E-103	1.0E-103	1.0E-103 A	1.0E-103	1.0E-103 X87831.2	1.0E-103 X87831.2	1.0E-103 AU136283 1	1.0E-103 L43610.1	1 00 400	1 05-103	1 0E-103 AB011300 1	1000	1 0F-104 AL	1 0F-104		1.0E-104 AA132975.1	4 OF 404 BEN44020.1	1.0E-104 BE	1.0E-104	1.0E-104	1.0E-104	1.0E-104 M34	1.0E-104 Y11151.1	1.0E-104 AA319436.1	1.0E-104 AB033102.1	1.0E-104 AB033102.1	1.0E-104 AB032998.1	1.0E-104 X02761.1
	Expression Signal	9.2	3.56	2.04	2.04	2.22	2.22	1.3	1.3	2.8	10.74	2 44	2.11	2 83	3.73	3.73	2.18	000	2.00	1 38	1.38	1.68	1.11	1.11	7.41	2.74	2.04	0.79	0.79	0.76	3.93
	ORF SEQ ID NO:	35998	36563							37340	36421	37667		31091	25878	25677	27328	27863	27688	27829	27830	27893	27965	27966	28285			28986	28987	29315	29712
	SEQ ID NO:	22786	23326	23424	23424	23434	23434	23476	23478	24037	23190	24339	24494	24620	13039	13039	14618	14919	14929	15091	15091	15158	15224	15224	15641	15683	16145	16341	16341	166/4	1/063
	Probe SEQ ID NO:	10138	10634	10737	10737	10748	10748	10783	10783	11347	11423	11748	11937	12128	227	227	1881	2190	2201	2369	2369	2438	2507	2507	2874	2917	3386	3587	_1	\$7AC	╛

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Table 4
Single Exon Probes Expressed in Brain

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Top Hit Descriptor	Homo sapiens chromosome 21 unknown mRNA	Homo saplens chromosome 21 unknown mRNA	Homo sapiens death receptor 6 (DR6), mRNA	Human Down Syndrome region of chromosome 21 DNA	Human Down Syndrome region of chromosome 21 DNA	Homo sapiens aik3 mRNA for Aurora/Ip11-related kinase 3, complete cds	wj03b12x1 NCI_CGAP_Kid12 Homo saplens cDNA clone IMAGE:2401727 3' similar to TR:Q14145 Q14145 KIAA0132 PROTEIN. ;contains element LTR7 repetitive element;	wi03b12.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2401727 3' similar to TR:Q14145 Q14145 KIA40132 PROTEIN :contains element LTR7 repetitive element	Homo saplens PDZ domain-containing guanine nucleotide exchange factor i (LOC51735), mRNA	601150451F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3503220 5'	601150451F1 NIH_MGC_19 Homo sepiens cDNA clone IMAGE:3503220 5'	Homo sapiens adaptor-related protein complex 2, beta 1 subunit (AP2B1), mRNA	UI-H-BI4-aow-b-09-0-UI:s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3086176 3'	nad16g11.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3365948 3'	zj98b05.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:462897 3'	yc83f02.r1 Soares Infant brain 1NIB Homo saplens cDNA done IMAGE:22440 5'			II.3-HT0619-080900-249-F07 HT0619 Homo saplens cDNA	IL3-H70619-080900-249-F07 H70619 Homo saplens cDNA	xd76d02x1 Sogres_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2603523 3' similar to TR:024116	miens cDNA clone IMAGE:2603523 3' similar to TR-O24116				601581503F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3935977 5'	AV728070 HTC Hamo saplens cDNA clane HTCBYA07 5'	AU130765 NT2RP3 Homo saplens cDNA clone NT2RP3001398 5'	24 and 25		RC0-HT0885-310700-021-b09 HT0885 Hamo saplens cDNA
Top Hit Database Source	NT	N	NT	N-	Ę	Z	EST_HUMAN	EST HUMAN	Ę	EST_HUMAN	EST HUMAN	Ā	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	MT	Nī	EST_HUMAN	EST_HUMAN	FOT LIMAN		EST_HUMAN	F	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	뉟	EST_HUMAN
Top Hit Acession No.	AF231920.1	AF231920.1	7657038 NT	U43379.1	U43379.1	AB017332.1	AI768797.1	AI768797.1	706512	BE314182.1	BE314182.1	11425572 NT	BF509244.1	BF448230.1	AA682308.1			AF091395.1	BF352841.1	BF352841.1	AW/103948 4		AW 103848.1	AF113514.1	BE791713.1	BE791713.1	AV728070.1	AU130765.1	U66535.1	27757	BE720191.1
Most Similar (Top) Hit BLAST E Value	1.0E-104		1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1 0E-104		1.0E-104		1.0E-104	1.0E-104		1.0E-104	1.0E-104		1.0E-104	1.0E-104	1.0E-104	1 OF 404		1.0E-104			1.0E-104	1.0E-104	1.0E-104		1.0E-104	1.0E-104
Expression Signal	1.23	1.23	1.06	1.26	1.26	1.12	9.51	9.51	0.75	3.31	3.31	2.03	0.83	5.23	0.5	1.31	4.27	4.27	4.4	4.4	נט		0.85	0.71	2.83	2.83	1.29	4.51	4.41	0.92	2.07
ORF SEQ ID NO:	29932			31573	31574		32141	32142	32322	32663	32664	32881	34330	34915	34997		35060	35061		02928	35503		35504	35698	35852	35853	36160	36204	36327		37221
Exon SEQ ID NO:	17305	17305		18637	18637	18682	19144	19144	1	19621	ı	19813	21187	21754	21833			21893	20412	20412	27306		22306			L	22946	22986	23096	23110	23930
Probe SEQ ID NO:	4570	4570	5106	5850	5850	5897	6375	6375	6551	90/9	6706	7125	8495	9065	9163	9184	9214	9214	9341	9341	0654		9654	9847	9993	9993	10299	10339	10450	10464	11268

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Top Hit Descriptor	RC0-HT0885-310700-021-b09 HT0885 Homo sapiens cDNA	602(41215F1 NIH_MGC_48 Homo saplens cDNA clone IMAGE:4302507 6'	601568806F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3841681 5'	801568806F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3841681 5'	Homo sapiens ribosomal protein S6 kinase, 90kD, polypeptide 5 (RPS6KA6), mRNA	601312181F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3658676 57	Homo sapiens amyloid beta (A4) precursor protein (protesse nextn-ll, Alzheimer disease) (APP), mRNA	Hamo sapiens Meis1 (mouse) homolog (MEIS1) mRNA	Homo saplens potasslum channel subunit (HERG-3) mRNA, complete cds	Homo sapiens potasslum channel subunit (HERG-3) mRNA, complete cds	Homo saplens chromosome 21 segment HS21C080	Human mRNA for KIAA0128 gene, partial cds	EST20609 Spieen I Homo sapiens cDNA 5' end similar to autoimmune antigen Ku, p70/p80 subunit	601434491F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3919511 5	Home saplens 959 kb contig between AML1 and CBR1 on chromosome 21q22; segment 1/3	Homo saplens bromodomain adjacent to zinc finger domain, 2B (BAZ2B), mRNA	Homo saplens bromodomain adjacent to zinc finger domain, 28 (BAZ2B), mKNA	EST373761 MAGE resequences, MAGG Homo captens cDNA	(801445823F1 NIH_MGC_63 Homo sapiens cDNA clone IMAGE:3850156 5	601445823F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3850156 5	Homo sapiens chromosome 21 segment HS21C008	Homo sapiens mRNA for KIAA0796 protein, partial cds	EST378088 MAGE resequences, MAGI Homo septens CUNA	Homo saplens E6-AP ubiquitin-protein ligase (UBE3A) gene, exon 2	Homo sapiens Retina-derived POU-domain factor-1 (RPF-1), mRNA	801901028F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4130334 5	601901028F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:4130334 5	Homo sapiens GTPase activating protein-like (GAPL), mRNA	Homo sapiens GTPase activating protein-like (GAPL), mRNA	EST363689 MAGE resequences, MAGB Homo sapiens cDNA	601677279F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3960019 5	Homo sapiens plakophilin 4 (PKP4), mRNA	Human mRNA for dbl proto-oncogene
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	TN	EST_HUMAN	L	NT	ΝΤ	LΝ	LN		EST_HUMAN	EST_HUMAN	LN	L	. 1	EST HUMAN		EST HUMAN	۲	NT	EST HUMAN	LZ LZ	ΙN	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	EST HUMAN	TN	ZT
Top Hit Acesslon No.	720191.1		1.0E-104 BE731976.1		11434729 NT	1.0E-104 BE393892.1	4502166 NT	4505150 NT	1.0E-105 AF032897.1	1.0E-105 AF032897.1	163280.2				229041.1	7304922 NT	7304922 NT			868881.1				1.0E-105 AF016704.1	11420134 NT			11419196 NT	11419198 NT	1.0E-105 AW951634.1	BE902616.1	6806894 NT	
Most Similar (Top) Hit BLAST E Value	1.0E-104 BE	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105 AL	1.0E-105 D50918.1	1.0E-105	1.0E-105	1.0E-105 AJ	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105 AL	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105 BE	1.0E-105	1.0E-105 X12556.1
Expression Signal	2.07	4.98	1.75	1.75	1.42	2.38	11.13	8.99	1.92	1.92	16.0	1.93	1.29	1.44	2.89	0.88	0.88	2.07	0.74	0.74	4.24	1.16	0.94	96'0	1.07	2.18	2.16	3,65	3.65	0.83	0.50	0.65	0.97
ORF SEQ ID NO:	37222	37259	37505	37508	37712		25722	25440	25987	25988	27269		27649			28759	28760		_				30428	30681		32485			30516				L
Exon SEQ ID NO:	23930	L	24189	24189	24381	l i	15514	12827	13360	13360	14554	L	1	1	١.	L	16105	١.	17428	17428	17613	17762	17810	18053	١.	1	L	Į.	١.	١.		1	1
Probe SEQ ID NO:	11288	11299	11590	11590	11791	12702	272	416	88	989	1814	1918	2186	2322	3008	3348	3346	4077	4694	4694	4886	5043	5091	5247	6312	6804	6804	6885	ARRE	6927	7184	7772	7758

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Single Exon Probes Expressed in Brain

7018c10.x1 NCI_CGAP_KId11 Homo sapiens cDNA clone iMAGE:3574291 3' similar to TR.P97880 P97880 Homo sapiens Ran binding protein 11 (LOC51194), mRNA w/4f07.x1 Soares_thymus_NHFTh Homo sapiens cDNA clone IMAGE:2535301 3' similar to TR:P87892 Human epidermai growth factor receptor (EGFR) precursor-mRNA, exon 4, partial ods ng41c05.s1 NCL_CGAP_Co3 Homo sapiens cDNA clone IMAGE:937352.3' similar to contains element ng41c05.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:837352 3' similar to contains element UHH-Blop-abi-b-12-0-UI.s1 NCI_CGAP_Sub2 Homo sapiens cDNA clone IMAGE:2711762 3 UI-HF-BN0-akt-g-07-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3078348 5 801443755F1 NIH_MGC_65 Homo seplens cDNA clone IMAGE:3847884 5' Homo seplens SMARCA4 isoform (SMARCA4) gene, complete cds, alternatively spliced EST02975 Fetal brain, Stratagene (cat#936206) Homo sapiens cDNA clone HFBCR32 ws50c10.x1 NCI_CGAP_Bm25 Hamo saplens cDNA clone IMAGE:2500626 3' similar Homo sapiens glutathlone S-transferase theia 1 (GSTT1), mRNA 801149783F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3502461 5' q176h10.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1878307 3' 601443755F1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3847884 5' tq79c01.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2215008 3 SW:ACSA_PENCH P36333 ACETYL-COENZYME A SYNTHETASE Homo sapiens glutathione S-transferase theta 1 (GSTT1), mRNA Fop Hit Descriptor RC1-CN0008-070100-011-e05 CN0008 Homo sapiens cDNA QV2-OT0062-140300-083-409 OT0062 Homo sepiens cDNA QV2-OT0062-140300-083-d09 OT0062 Homo sapiens cDNA EST377629 MAGE resequences, MAGI Homo sapiens cDNA MR0-HT0165-140200-008-d10 HT0165 Homo sapiens cDNA Homo sapiens COL4A6 gene for a6(IV) collagen, expn 31 Homo sapiens soluble neuropilin-1 mRNA, complete cds Human dihydrofolate reductase pseudogene (psi-hd1) Human dihydrofolate reductase pseudogene (psi-hd1) Homo sapiens gene for Smed 3, exon 2 and 3 Homo sapiens gene for Smad 3, exon 2 and 3 LTR3 repetitive element; LTR3 repetitive element P87892 PROTEASE RIN. EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST HUMAN HUMAN EST_HUMAN HUMAN Top Hit Database Source EST_HUMAN HUMAN EST_HUMAN ES F RIST 눋 눋 7705936 NT È 4504184|NT 4504184 NT Top Hit Acession 1.0E-105 AW007194.1 1.0E-105 AW840817.1 1.0E-105 AW016879.1 1.0E-105 AW882372.1 1.0E-105 AW882372.1 1.0E-105 BE867793.1 1.0E-105 AW027554.1 1.0E-105 AB004924.1 1.0E-106 AW 503208.1 1.0E-108 AI565065.1 1.0E-108 AW965556.1 1.0E-108 J00146.1 1.0E-106 BE260201.1 1.0E-106 AI276528.1 1.0E-105 BE867793.1 1.0E-105 BF430921.1 1.0E-105 AB004924.1 AA527446.1 1.0E-106 AA527446.1 1.0E-106 BE144286.1 ģ AF145712. 05087.1 1.0E-108 U48724.1 D83548. 1.0E-106 J00146.1 1.0E-106 1.0E-105 0E-106 .0E-105 1.0E-105 1.0E-106 .0E-106 (Top) Hit BLAST E fost Simila 2.82 1.73 1.39 2.68 0.75 6.0 2.06 3.48 6.55 99.0 0.68 5.73 1.59 2.58 1.73 99.0 4.7 4.71 2.31 1.82 3.24 1.91 Expression Signal 35247 35248 34786 34954 27246 27578 27778 28064 28210 33750 34955 36776 37494 37592 25654 28000 26000 27138 27245 26842 ORF SEQ ΩNÖ 24179 13013 20622 23860 SEQ ID 21791 23912 24270 24415 12962 13313 13371 13371 14262 14440 14538 14536 15040 15322 15466 14159 Š ö 9103 SEQ ID 8297 8828 8948 9103 9467 11580 11675 2118 2810 9467 11195 200 529 591 1515 1796 2828 11250 1831 147 592 7927 10850 11831 1798 1697 ÿ

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Single Exon Probes Expressed in Brain	Top Hit Descriptor		Tromo suprems grutatritione S-transferase theta 1 (GSTT1), mRNA	nomo sapiens mKNA tor KIAA1326 protein, partial cds	nomo sapiens mknA for KIAA1328 protein, partial cds	Inomo sapiens hypothetical protein FLJ11273 (FLJ11273), mRNA	Indino sapiens hypothetical protein FLJ11273 (FLJ11273), mRNA	Homo sapiens gene for activin receptor type IIB, complete cds	Induit suprems mixINA for KIAA1278 protein, partial ods	Homo sapiens mKNA for KIAA1278 protein, partial cds	ESTSSSST MAGE resequences, MAGN Homo sapiens cDNA	Home services April 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	MBD-HT0188 440200 000 420 110 000 000 000 000 000 000 000 000 0	H. sapiens mRND similar to D29763 mouse mRNA for selzure-related gene product 6. Shares domains with	H. saplens mRNA similar to D29763 mouse mRNA for seizure-related gene product 6. Sheres domeins with	BMIPs, I olloid, Sushi repeat proteins al24b09.s/ Sorres testis NUT Home	PHPS1-2 (HUMAN);	AU130113 NT2RP3 Homo saplens cDNA clone NT2RP3000274 F	AU130113 NT2RP3 Homo sapiens cDNA clone NT2RP3000274 8'	zw28d12.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:770815.31	AU143428 Y/8AA1 Homo saplens oDNA clone Y79AA1001912 5	AD 143442 778AA1 Homo saplens cDNA clone Y79AA1001912 6	601439620E4 NILL MGC 3211	Home Scales Control Machine Control Representations IMAGE:3924841 5'	Homo saplens Aylosytransferese II (XT2), mRNA	ae72607 s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:969737 3' similar to chi-Yazaza	KINESIN HEAVY CHAIN (HUMAN);	Homo saplens XPIMC2 protein (LOC57109), mRNA	OVI 1037 39F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE: 2988345 61	Home suprems sorting nexts 11 (SNX11), mRNA	AU118850 HEMBA1 Homo sapiens oDNA clone HEMBA1000129 F	D RZI DODI VOLICIO I COLOR I C
Igle Exon Pro	Top Hit Database Source	Ę	L			L L	LV	L L	F	FOT LIMAN	FOT HIMAN	NUMBER OF THE PROPERTY OF THE	T HI MAN	T^{-}					П	EST HUMAN	Т	Т	Т	Т			NAMOL	T III IMAANI	NEW PARTY		T_HUMAN	
	Top Hit Acession No.	4504484 NT	AB03774	1.0E-106 AB037747 1	TO SOCK AIT		ABOORES	E-106 AB033104 1	-106 AB0331041		ı] <u>č</u>	1.0E-106 BE144286.1							100 AA434106.1	T	T	Γ	15913	11545913 NT	108 44683770 4	20847	106 BE292722 1 E	5503	11425503 NT	106 AU116850.1	
	Most Similar (Top) Hit BLAST E Value	1.0E-108	1.0E-106	1.0E-106	1.0E-108	1.0E-108	1.0E-10B	1.0E-106	1.0E-106	1.0E-108	1.0E-106	1.0E-106	1.0E-108	1.0E-106	1.0F_108		1.0E-106	1.0E-106/	1.001-100	1 0F-108 /	1.0E-108 A	1.0E-106B	1.0E-106 B	1.0E-106	1.0E-106	1 0F-108					1.0E-106 AI	
	Expression Signal	1.91	5.23	6.23	2.18	2.18	0.81	1.18	1.18	7.95	7.95	1.05	1.4	1.09	1.09		2.61	0.58	000	13	1.3	4.89	0.77	17.66	17.66	5.16	5.33	1.21	9.29	9.29	0.67	
	ORF SEQ ID NO:	26843	28357	28358	28588	28589	28784	28838	28839	29391	28392	29410	29924	30469	30470		30750	31481	31533	31637	31638	31755	31860	32083	32084	33031	33084	33161	33275	33276	33491	
	<u> </u>			i I	15939	1 1	16126	16190	16190	16763	16763	16780	17297	17853	17853	0000,	18090	18555	18805	18889	18689	18792	18892	19095	19095	19955	20002	20080	20182	20182	20378	
	Probe SEQ ID NO:	2828	2942	2942	3176	3176	3367	3434	3434	4017	4017	\$ 233	4562	5135	5135	2002	5764	5764	5816	5904	5904	6011	8303	8325	6326	7271	7324		1	Ш	7714	

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	Top Hit Descriptor	EARBOARATE1 NIH MGC 9 Homo sapiens cDNA clone IMAGE:3948463 5'	6015943311 1111 MGC 9 Homo saplens cDNA clone IMAGE:3948463 5	or 82 n.7 v. Parstead Borta HPLRB6 Homo sapiens cDNA clone IMAGE:2127732 3' similar to gb:X06233	CALGRANULIN B (HUMAN);	601282717F1 NIH_MGC_44 Homo sapiens cDNA cigne INAACE-5504403 R	601282717F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE.Southess 3	NG2B05.X1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE: 2260362 5 SIIIIIII COMP Kid11 Homo sapiens cDNA clone IMAGE: 2260362 5 SIIIIIIII COMP	Q05084 69 KD ISLET CELL AUTOANTIGEN ;	CARTAINS INC. CGAP GCB1 Homo saplens cDNA clone IMAGE:1354790 3'	ACAZANS 31 NCI CGAP GCB1 Homo sapiens cDNA clone IMAGE:1354790 3	Helphare PC-CC CCLI III	cn03a04.71 Normal Human Trabecular Bone Cells Homo saplens cDNA clone NH i BC_cn03a04.41 millouning to contains MSR1.13	ITAR1 PTR5 repetitive element;	Ima1102 x1 NCI CGAP Kid11 Home saplens cDNA clone IMAGE:2160699 3' similar to contains Mont. La	TAR1 PTR5 repetitive element :	80 (262507) 1111 MGC 20 Homo saplens cDNA clone IMAGE:3954403 5	PATA PATA NIH MGC 20 Homo sepiens cDNA clone IMAGE:3954403 5	BOTO 10.1 NOI CGAP BI2 Homo sepiens cDNA clone IMAGE:1130395 3	INDS7b10.81 NCI CGAP_Br2 Homo sepiens cDNA clone IMAGE:1130395 3	RC0-CT0318-201199-031-a11 CT0318 Homo saplens cDNA	DKFZp434F0712_11 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434F0712.5	Homo sapiens chromosome 21 segment HSZ1 C002	601453461F1 NIH MGC_68 Homo sapiens GDNA clone IMAGE: 3857368 5	601453461F1 NIH MGC 66 Homo sapiens curvo con a monoconoconoconoconoconoconoconoconocono	Human ryanodine receptor mRNA, complete cds	Human ryanodine receptor mRNA, complete cds	601109219F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3345397 3	RC5-BN0192-100500-021-B02 BN0192 Homo sapiens cDNA	RC5-BN0192-100500-021-B02 BN0192 Homo saplens cDNA	fh05h11x1 NIH MGC_17 Homo sapiens cDNA clone IMAGE:2961644 5	601433087F1 NIH_MGC_72 Hamo sapiens cDNA clone IMAGE:3918524 5	
- I SVI OBIIIO	Top Hit Database Source		EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN		EST HUMAN	TOT TOT	TOT TOWAR	בא החשואות	EST_HUMAN	EST HIMAN	NUMPI I SE	EST_HUMAN	EST HUMAN	ESI DUMAN	TOT HOMAN	EST HUMAN	EST HIMAN	EST HUMAN	N	EST_HUMAN	EST HUMAN		\ V	FST HUMAN	EST HUMAN	NAM! H TOO	NAMINAAN	EST HUMAN	
5	Top Hit Acession No.		BE741408.1	BE741408.1	A1523066.1	RF387950.1	BE387950 1		A1654123.1		AA825307.1	AA825307.1	AI750447.1		A1479509.1			_	S BF027310.1		5 AA604417.1	5 AW 303239.1		BF032755.1	B BF032755.1	6 In5200 1	1.0E-1001300200.1	1.0E-100,000,00.1	0 DECA1300.1	1.0E-106 BE010802.1	1.0E-106 BE010882.1	1.0E-106 AW 410403.1	
	Most Similar (Top) Hit BLAST E	A dina	1.0E-106	1.0E-106	1 0F-108	4 0E-108	10.1	201-120	1.0E-106	1.0E-106		1.0E-106	1.0E-106		1.0E-106							1.0E-105	\perp	┸	1	405			1				_[_
	Expression Signal		6.44	6.44	1 38	25.0	0.47	***	3.64	0.58	3.28	3.28	1.28		1.8	1.8	1.19	1.09	1.09		·			707									3.32
	ORF SEQ ID NO:		33707	33708	20000	١		34362	34443		34889	34890	35021		35169	35170	<u> </u>	35841								1					37437		31068
	SEQ ID	<u> </u>	20579	20579		20768	21219	21219	21299	21641	l .	1	21856	1	21996	21996	1	L	1	1_	<u> </u>	1)		- 1	1				4 23991	2 24132	1	L	3 24664
,	Probe SEQ ID	į	7884	7884		8074	8527	8527	8607	8950	9045	9045	98.60	3	9329	9329	9902	8084	988	10141	10141	10185	10358	10479	10812	10812	10995	10995	11384	11532	11532	11981	12193

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VO	01/57275	_	Т	Ť	Ť	Т	Ť	_	Т		_	7	1	T	1	_	_	_	_	, '	-	¥	, 1	_		·	P <u>C</u>	<u> </u>	i <u>S</u> 0	1/0)06	67
Single Exon Probes Expressed in Brain	Top Hit Descriptor		601433087F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3918624 5'	RC1-CT0249-090800-024-d05 CT0249 Homo sapiens cDNA	Homo sapiens Xq pseudoautosomal region; segment 1/2	Human IFNAR gene for interferon alpha/beta receptor	Homo sapiens neuronal cell adhesion molecule (NRCAM) mRNA	Inomo sepiens NY-KEN-25 antigen mRNA, partial cds Human IENAR nans for Interference IIII in the constant of the constant in th	The second application application applications are second as the second application are second as the second as t	Homo sepiens sodium-dependent high-affinity dicarboxylate transporter (NADC3) mRNA, complete ode	Homo sepiens BAZ1B mRNA for bromodomain adjacent to zinc finger domain 1B, complete ods	4V2-H10540-120900-358-a05 H10540 Homo saplens cDNA	nomo sapiens cathepsin Z precursor (CTSZ) gene, exon 3	Homo sabiens mRNA for KIAA0453 protein, partial cds	Homo sapiens mRNA for KIAA0453 protein, partial cds	Plan Palesta (2015) gene, exen 20	DIM CHOOSE 190100-001-003 CN0031 Homo sapiens cDNA	ENT-CINUS I-19010U-001-003 CN0031 Homo sapiens oDNA	SOLOCIO SEL INIT., MCC., 21 Homo saplens cDNA clone IMAGE:3842309 5	DAM CALORA 400,000 001 1000 Saplens CDNA clone IMAGE:3842308 67	PA41-CANDRA 4 400000 ART 100 COURT Homo saplens cDNA	Hamperson Control of the Control of	Home septents own I 3 (suppressor of mif two 3, yeast) homolog 2 (SMT3H2), mRNA	FETTER 14 FLANCE	601442558E1 NIH MGC 65 Home capiens conv	September 201 John September 2014 Globe IMAGE:3848484 6	norto septens general transcription factor IIIC, polypeptide 1 (alpha subunit, 220kD) (GTF3C1), mRNA	Homo Saplens general transcription factor IIIC, polypeptide 1 (alpha subunit, 220kD) (GTF3C1) mRNA	U-HT-BNI0-aff-c-08-0-UI.1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3078310 6'	UFTIT-BING-BIT-C-08-0-ULT NIH MGC 50 Homo saplens cDNA clone IMAGE:3079310 5'	#INSOID=XI NC_CCAP_Kid11 Homo sapiens cDNA done IMAGE:2384791 3'	AU122469 MAMMA1 Homo septems cDNA clone IMAGE:2964524 5'
igle Exon Pr	Top Hit Database Source	1	HOMAN	EST_HUMAN	N.	Z		I Z	<u> </u>	1	ROT CIRAN	HA LES	H	H	Z	EST LIMANI	EST HIMAN	EST LIMAN	EST HIMAN	EST HIMAN	EST HIMAN	L.	Į.	FRT HIMAN	EST HUMAN				EST HIMAN	7	7	Т
ES	Top Hit Acession No.	DE004400 4	BEGS4400.1	A.1274735 4	X60450 4	4826863 NT	AF155103 1	T	AF154121 1	T	T	T	I	T	T		T		T	Ī	T	2002	F020671.1		T	71434480 NIT		ĝ	T	T	T	
	Most Similar (Top) Hit BLAST E Value	1 OF 10R	1.0F-10R	1.0E-107	1 0F-107	1.0E-107		_	1.05~107		_	-	_	-		+	1.0E-107	_	-	-	1~	1.0E-107	1.0E-107 A	1.0E-107 A	1.0E-107 B	1.05-107	1 OF 107	1 0E-107 AWE03042 4	1.0E-107/AW503913 1	1.0E-107 AI	1.0E-107 AW 410961.1	1.0E-107 AU122469.1
	Expression Signal	3.32	4.6	4.42	18	4.03	1.88	1.62	11.55	0.73	3.77	1.56	0.99	0.99	0.95	1.45	1.45	1.2	1.2	3.89	3.89	2.63	5.14	99'0	3.2	1.45	1 45	1.42	1.42	1.48	9.0	0.95
	ORF SEQ ID NO:	31069					26028	26305	26379	26673	26986	27187	27283	27284	27871	27822	27823	27891	27992	28421	28422	28602	29190	31242	31495	32508	32507	33023	33024	33183	33469	35213
	SEQ ID	24664		L	L	1 1	13394	13635	13714	14004	14308	14488	14871	14571	14933	15084	15084	15250	15250	16773	16773	15861	18558	18335	18566	19484	19484	19947	19947	20106	20354	22041
	Probe SEQ ID NO:	12193	12408	228	258	607	616	998	948	1266	1562	1746	1832	1832	2205	2362	2362	2635	2635	3007	3007	3096	3806	5537	57.75	6823	6823	乚	L			8287

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Table 4
Single Exon Probes Expressed in Brain

Top Hit Descriptor	tg10d08.x1 NCI_CGAP_CLL1 Homo saplens cDNA clone IMAGE:2108383 3' similar to SW:AACT_DICDI P05095 ALPHA-ACTININ 3, NON MUSCULAR ;	Homo sapiens neuroendocrine-specific protein (NSP) gene, exon 4	602123963F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4281039 5'	601066681F1 NIH_MGC_10 Homo sapiens cDNA clane IMAGE:3452829 5	Homo sapiens HSPC049 protein (HSPC049), mRNA	Homo sapiens HSPC049 protein (HSPC049), mRNA	Homo sapiens solute carrier family 10 (sodium/bile acid cotransporter family), member 1 (SLC10A1) mRNA	Homo sepiens solute cerrier family 10 (sodium/bile acid cotransporter family), member 1 (SLC10A1) mRNA	ze45e01.s1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:361944 3' similar to contains THR.b1 THR repetitive element ;	601177018F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3532348 5	Homo saplens NF2 gene	bb25b10x1 NIH_MGC_14 Homo sapiens cDNA cione IMAGE:2963899 3' similar to gb:X63777 60S RIBOSOMAL PROTEIN L23 (HUMAN); gb:J05277 Mouse hexokinase mRNA, complete cds (MOUSE);	Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds	Homo saptens potassium channel subunit (HERG-3) mRNA, complete cds	hitzart.xr NCI_CGAP_GU1 Homo sapiens cDNA clone IMAGE:2972060 3' similar to SW:3BP1_MOUSE_ P55194 SH3-BINDING PROTEIN 3BP-1.;	Human hepatocyte nuclear factor 4-alpha gene, exon 2	Human hepatooyle nuclear factor 4-alpha gene, exon 2	Homo sapiens KIAA0187 gene product (KIAA0187), mRNA	Homo sapiens PSN1 gene, alternative transcript	RCO-HT0372-241189-031-d03 HT0372 Homo sepiens cDNA	601444922F1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3848980 5'	601444922F1 NIH_MGC_65 Hamo sapiens cDNA clone IMAGE:3848980 5'	Homo sapiens familiai mental retardation protein 2 (FMR2) gene, exon 20	Homo sepiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2 mRNA, complete cds	Homo sapiens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2 mRNA, complete cds	
Top Hit Database Source	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN					EST HUMAN	EST HUMAN	П	EST_HUMAN	Г	LN	EST_HUMAN	Į.	- LZ		LN	EST_HUMAN	EST_HUMAN (Г		Į.	N TN	
Top Hit Acession No.	1.0E-107 Al392850.1	-107 L49141.1	-107 BF666511.1	1.0E-107 BE540550.1	11419701 NT	11419701 NT	4506970 NT	4506970 NT	-107 AA001415.1	-108 BE296042.1		-108 BE206694.1		-108 AF032897.1	-108 AW664438.1	-108 U72961.1	-108 U72961.1	7661979 NT	-108 AJ008005.1	108 AW384094.1	-108 BE869016.1	1.0E-108 BE869016.1	108 AF012623.1	-108 AF264717.1	108 AF264717.1	
Most Similar (Top) Hit BLAST E Value	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-107	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	
Expression Signal	3.18	14.26	1.71	99.9	5.97	5.97	1.36	1.36	5.86	2.66	1.87	4.97	0.71	0.71	1.14	2.18	2.18	1.74	2.8	1.15	1.7	1.7	66.0	6.13	6.13	
ORF SEQ ID NO:	36537		36807	37252			37768	37769			26657			28756				30113	30241	30883	30954	30825		31790	31781	
SEQ ID NO:	1	23546	1	L	23178	23178	24427	24427	25328	13702	il		16103	16103	16878		17225	17484	17623	18191	18239	18239	18626	18828	18828	
Probe SEQ ID NO:	10603	10866	10880	11293	11367	11367	11843	11843	12043	835	1242	2428	3344	3344	4136	4489	4489	4752	4886	5391	5440	5440	5837	6048	6048	

Page 438 of 536 Table 4 Single Exon Probes Expressed in Brain

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Top Hit Descriptor	Homo eapiens caveolin-1/-2 locus, Contig1, D7S622, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and 2)	PM4-CT0403-240700-001-c10 CT0403 Homo saplens cDNA	Homo sapiens E6-AP ubiquitin-protein ligase (UBE3A) gene, exon 4	Homo sapiens E6.AP ubiquitin-protein ilgase (UBE3A) gene, exon 4	Homo saplens G protein-coupled receptor, family C, group 5, member B (GPRC5B), mRNA	Homo saplens delta-6 fatty acid desaturase (FADSD8) mRNA	601113471F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3354064 5'	602043384F1 NC _CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4181037 5'	602043384F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4181037 5'	Homo saplens connective tissue growth factor-like protein precursor, mRNA, complete cds	UI-HF-BM0-ads-e-12-0-UI.r1 NIH_MGC_38 Homo sapiens cDNA clone IMAGE:3082878 5	UI-HF-BM0-ads-e-12-0-UI.r1 NIH_MGC_38 Homo sapiens cDNA clone IMAGE:3062878 5	Homo sapiens ETS-family transcription factor EHF (EHF) mRNA, complete cds	1y35h10.r1 Soares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:273283 5' similar to PIR:A48773	A45773 kelch protein, lang farm - fruit fly;	601058769F1 NIH_MGC_10 Homo saplens cDNA clone IMAGE:3445361 6'	Homo sapiens mRNA for Golgi-associated microtubule-binding protein (GMAP-210)	Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternativaly	spliced	EST378268 MAGE resequences, MAGI Hamo sapiens cDNA	AV708790 ADC Hamo saplens cDNA clane ADCAEE03 5'	AV708790 ADC Homo saplens cDNA clone ADCAEE03 5'	Homo sapiens G protein-coupled receptor 48 (GPR48), mRNA	Homo sapiens COL4A6 gene for a6(IV) collagen, exon 23	Homo sapiens mRNA for FLJ00037 protein, partial cds	602018571F1 NC _CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4154297 5	IL2-UM0077-260400-079-D06 UM0077 Homo saplens cDNA	Human mRNA for KIAA0220 gene, partial cds	Homo sapiens reticulocalbin 1, EF-hand calcium binding domain (RCN1), mRNA	Homo saplens tetratricopeptide repeat domain 2 (TTC2) mRNA	Homo saplens mRNA for KIAA0899 protein, partial cds	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds
Top Hit Database Source	NT	EST_HUMAN	LΝ	١	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝΤ	EST_HUMAN	EST_HUMAN	ΙN		EST_HUMAN	EST_HUMAN	攴		ΝŢ	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN.	NT	±Ν	EST_HUMAN	EST_HUMAN	Į.	F	۲	ĻΝ	NT	۲
Top Hit Acessian No.	AJ133269.1	1,0E-108 BF334851.1	1.0E-108 AF016706.1	1.0E-108 AF016708.1	11431857 NT	475833 NT	1.0E-108 BE252607.1	1.0E-108 BF528912.1	1.0E-108 BF528912.1	1.0E-108 AF083500.1	1.0E-108 AW 408694.1	1.0E-108 AW 408694.1	1.0E-108 AF203977.1		1.0E-108 N44974.1	BE635227.1	1.0E-108 Y12490.1		1.0E-108 AF223391.1	1.0E-108 AW966185.1	1.0E-108 AV708790.1	AV708790.1	11441465 NT	D63539.1	AK024447.1	1.0E-108 BF346356.1	1.0E-109 AW803116.1	D86974.1	11438391 NT	4507712 NT	1.0E-109 AB023216.1	1.0E-109 AB023216.1	M28699.1
Most Similar (Top) Hit BLAST E Value	1.0E-108 A.	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108		1.0E-108	1.0E-108	1.0E-108		1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108 D	1.0E-108 A	1.0E-108	1.0E-109	1.0E-109 D86974.1	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109
Expression Signal	1.27	0.92	0.63	0.63	5.82	3.55	1.16	0.84	0.84	1.68	0.48	0.48	0.75		0.54	1.73	1.98		1.39	3.82	2.2	2.2	1.67	1.68	2.41	8.32	78.0	76.0	1.59	5.59	26.8	26.8	10.97
ORF SEQ ID NO:	31918	31650	32290	32291	32811	33098	33137	33164	33165		33838	33839	34781		34823	33481	30597		36928	37191		37254		37369	31072		25488	25523	25667	25878	25992	25993	26598
Exen SEQ ID NO:	18946		L	19287	19748			20082	20082	20638	乚	L.	21636	L	21674	20368	L		23671	23902		23955	24033	24054	24671	L	1_	L	1	Ì.		13364	13833
Probe SEQ ID NO:	6169	6265	6521	9521	7057	7339	7377	7405	7405	7963	8014	8014	8945		8984	10565	10731		10998	11239	11294	11294	11343	11405	12204	12583	41	62	220	454	288	584	1180

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Most Similar Top Hit Acession Database BLAST E No. Source	LN	1.0E-109 BE283673.1 EST_HUMAN 601186922F2 NIH_MGC_15 Homo sapiens dJNA done inva-ca-sacco o	EST_HUMAN	N	1.0E-109 AL163284.2 NT	1.0E-109 Y17123.1 NT		1.0E-109 AI022328.1 EST_HUMAN	1.0E-109	1.0E-109 N85190.1 EST_HUMAN	1.0E-109 AW893192.1 EST_HUMAN	1.0E-109 AW893192.1 EST_HUMAN	1.0E-109 AF240698.1 NT	1.0E-109 M37928.1 NT	1.0E-109 M37928.1 NT	1.0E-109 BE146144.1 EST HUMAN	1.0E-109 AB011181.2	1.0E-109 AB011181.2 NT	1.0E-109 AI655417.1 EST_HUMAN	1.0E-109 AA682274.1 EST_HUMAN	1.0E-109 AA662274.1 EST_HUMAN	1.0E-109 4504206 NT	1.0E-109 7662083	1.0E-109 R15400.1 EST HUMAN	1.0E-109 BE293673.1 EST HUMAN	1.0E-109 BE283673.1 EST HUMAN	1.0E-109 AU13/282.1 EST HIMAN	1.0E-109 BF6/3/18.1
							17	1.0E-109 Al022328.1							·				09 AI655417.1	1.0E-109 AA682274.1	09 AA66227			1.0E-109 R15400.1	1.0E-109 BE293673.1	1.0E-109 BE293673.1	1.0E-109 AU13/282.1	1.0E-109 Bro/3/10.1
Expression (To Signal BL	4			L	L					2.22	3.14	3.14	121			2.59	1.42	1.42	3.88			2.48						1.06
ORF SEQ ID NO:	SOROR	26067	28968									İ					28357		29497				\		30320	П		30503
Exan SEQ ID NO:	42022		1	L	-1	1	1	l	١.	<u> </u>	┸	L	L	1	i_	L	1	1	16869	1	<u> </u>	L	L	┺	Ĺ	17715	1	17988
Probe SEC ID NO:	7077	0 2	32	230,	1807	2248	2628	2828	2629	2054	5000	3383	3508	3548	3548	3823	3974	3974	4127	4147	4141	4371	458	4867	4992	4992	5167	5179

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WO 01/57275

				Γ	Г	Г	Г	Г	Т	Γ		_	г	Т	Γ-		$\overline{}$	_	т	•		-	•	写得	Ŧ;	=,	_,	<u>~</u>	- 2	-	4		<u>ه ب</u>	-
Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sealone njevnetnje 44 / - 1	RC1-HT0815-200400-022-404 Ltzsstr. 1	CM1. H70038-080006-202-004 H10816 Homo sapiens cDNA	PC4. HT0845. 200406. 2022 104. LT0825.	Home conference and A 1773 A 1	Homo septens minner for NIAA1691 protein, partial ods	SOURCES A Family United the Source of A 18F1), mRNA	SOURCESSON IN THE WIND TO JOIN SEPTEMBERS CLUNA CIONE IMAGE: 4040279 5'	801145017F2 NIH MCC 10 Home Saplens cUNA clone IMAGE: 4040279 5'	Novel Triman dens manalita to the control of the co	SACE THIRM BOTH MAPPING TO CHOMOSOMB 13	7818HING Chromoson 7 F-41 Butter appeals conversed to the converse conversed to the converse conversed to the converse conversed to the converse co	SO147047754 NILL MCC 6911	601479447F1 NIH MCC 69 H	II 0-HT0205-071100-142-44 HT0205 U	ys90g08.11 Sogres retina N2b6HR Homo septens cDNA clone IMAGE: 222110 6' similar to SP-A53491	A53491 BUME I ANIDE-SENSITIVE NA-K-C1 COTRANSPORTER - SPINY;	CONTROL OF THE WIND WIND BEAUTION OF THE WAR CONTROL OF THE WAR CONTROL OF THE WIND WIND SET THE WAR CONTROL OF THE WAY CONTROL	HSCHEC121 pormelized infant built. April 11	601063030F1 NIH MGC 10 Homo conjunt ANN ANN ANN ANN ANN ANN ANN ANN ANN AN	601063030F1 NIH MGC 10 Home Bahlens china Alexan MA CE 3449599 6	602080724F2 NIH MGC 81 Homo saniens conta clara (NA OF 1212111)	Homo saplens KIAA0744 dene product: historie descentars 27/kia A 274/3	Homo sapiens KIAA0744 cene product historie description 7 (214 2074)	AU121370 HEMBB1 Home seniens CONA class HEMBB100200 FT	Homo saplens Chedlak-Higashi swidnima 1 (CHS1) meNA	화8612.r1 Soares_feta[_lung_NbHL19W Homo saplens cDNA clone IMAGE:301439 6' sImilar to	Homo sepiens single-minded (December 1971)	BODO39003E1 NCT CCAB Bross U.	602039003E1 NCI_CGAP_Brist Home sablens cUNA clone IMAGE:4188753 5'	Homo sapiens SNF5/INIT gene even 8	Homo seplens SNF6/INI1 gene, exon 6	
gle Exon Prot	Top Hit Database Source	L	EST HIMAN	EST HIMAN	Т	Т		H HIMAN	Т	Т	Т	H IMAN	Т	Т	Т	Т	Т	EST HUMAN	Т	Т	Т	Т	EST HUMAN 6			EST HUMAN A		EST HIMAN		T HUMAN	T	Ĭ		
Sin	Top Hit Acession No.	6174822 NT	BE17835	109 BF379688.1	-109 BE179358 1	T	2574	109 BF182707.1		T	Γ		T	T	Γ	Π	000 L94060 4	-	T		-	09 BE540909.1 E	09 BF694831.1 E	7662279 NT	7662279 NT	109 AU121370.1 E	838	09 W16510.1	418618		l			
	Most Similar (Top) Hit BLAST E Value	1.0E-109	1.0E-109	1.0E-109		1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1 0E-100		1.0E-109	1.0E-109 F	1.0E-109 B	1.0E-109 E	1.0E-109 B	1.0E-109	1.0E-109	1.0E-109 A	1.0E-109	1.0E-109 W	1.0E-109	1.0E-109 BF339540.1	1.0E-109 BF339540.1	1.0E-109 Y17123.1	1.0E-109 Y	
	Expression Signal	3.09	1.11	0.64	1.6	0.97	3.89	5.28	6.28	0.67	1.48	0.89	2.77	8.42	8.42	0.56	- 6	0.63	0.63	2.64	1.71	1.71	15.79	1.65	1.55	1.8	2.72	11.6	1.46	1.27	1.27	2.1	2.73	
	ORF SEQ ID NO:	30662		31560		32896	33230	33232	33233	33461	33896	34014		34466	34467	34711	34990	35101	35102	35218	36606	36607	36638	36818	36819	37013	37338	37291	37602	37773	37774	27714	27714	
	SEQ ID	L						20140		' I	- 1	ı			_ I	21567	21825	21929	21929	22046	23364	23364	23399	23068	23568	23739	24032	23990	24280	24432	24432	14976	14976	
	Probe SEQ ID NO:	5230	2520	6838	5907	7140	7464	7466	7466	7674	873	8183	8665	8633	8833	8878	9137	9250	9250	9384	10673	10673	20/12	2000	2000	11069	11342	11383	11685	11848	11848	12112	12328	
٠																										_			_ •			_'.		

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Most Similar Expression (Top) Hit Top Hit Acession Signal BLASTE No. Source	2.08 1.0E-109 AB01:138	1.65 1.0E-110 7549804 NT	4.71 1.0E-110 5803073 NI	4.71 1.0E-110 5803073 NT	0.7 1.0E-110 C04498.1 EST_HUMAN	2.26 1.0E-110 7549804 NT	1.54 1.0E-110 U84550.1 INT	0.8 1.0E-110 5031620 N1	0.8 1.0E-110 AB032263.1 N1	1.19 1.0E-110 BE379477.1 EST HUMAN	1.6 1.0E-110 BF508896.1 EST HUMAN	4503098 NT	1.49 1.0E-110 11436041 NT	1.49 1.0E-110 11436041 NT Homo sapiens pregnancy_train privers (1.4.5), iii. v. 7, iii.	NAMUH TRE	1.09 1.05-110 050 1030.1	2.14 1.0E-110 AI017213.1 EST_HUMAN	3.9 1.0E-110 AU117812.1 EST HUMAN	2.7 1.0E-110 7662441	2.63 1.0E-110 BE289408.1 EST HUMAN	0.8 1.0E-110 BE621069.1 EST_HUMAN	8.61 1.0E-110 11419323 NI	8.61 1.0E-110 11419323 N1	8.08 1.0E-110 M55112.1 NI	0.8 1.0E-110 U08888.1 INI	0.8 1.0E-110 U08888.1 NI	0.74 1.0E-110 AIS60289.1 EST_HUMAN	6.9 1.0E-110 AV714276.1 EST_HUMAN	6.9 1.0E-110 AV714276.1 EST HUMAN	3.21 1.0E-110 AB020875.1	0.83 1.0E-110/AU137923.1 [EST_HUMAN
				Ц								1 1.0				1														Ш	Ц
Exen ORF SEQ E SEQ ID NO:	24813 31047	12831 25444	12864 25482	12864 25483	L			13912 26575	14005 26674			15613	15952 28603	L]_	16776 29407	17328 28951		L	18020 30642		18451 31384	18451 31365	25096 32395	19694 32748	19694 32747	19909	L			
Probe SEQ ID	12443	1	36	<u>l_</u>	62	107	514	1157	1256	1914	2051	2845	3189	3189		4031	7.04	4808	4916	5212	5639	2656	5656	6617	7002	7002	7007	7325	7325	7355	7469

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Probe SEO ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9234	21913	35087	7.88	1.0E-110	O BE302594.1	EST_HUMAN	ba68f01.y1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2805561 5' similar to TR:077258 077258 EG:114D9.2 PROTEIN.;
9478	L		2.39	1.0E-110	10 AW838394.1	EST HUMAN	QV2-LT0063-020400-119-e04 LT0053 Homo sapiens cDNA
10221	22869	36081	3.45	1.0E-110	11432732 NT	Z	Homo sapiens galactokinase 2 (GALK2), mRNA
10648	23339	36578	3.64	1.0E-110	10 Y12337.1		H.saplens mRNA for myotonic dystrophy protein kinase like protein
10887			3.75	1.0E-110		EST_HUMAN	601565604F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3840433 5
10887		36817	3.75	1.0E-110	10 BE734357.1	EST_HUMAN	601565604F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3840433 5
11420	<u></u> _	36418	2.45	1.0E-110	10 AA446529.1	EST HUMAN	zw67g02.r1 Soares_tests_NHT Homo saptens cDNA clone IMAGE:781288 5' similar to TR:G1145816 G1145816 FKBP54 ;
11939	1_			1.0E-110		EST_HUMAN	601439784F1 NIH_MGC_72 Homo saplens cDNA clane IMAGE:3824548 5'
12061	┸		11.71	1.0E-110	10 AW062258.1	EST_HUMAN	IL0-BT0163-040899-094-g10 BT0163 Homo seplens cDNA
12290			4.5	1.0E-110	1.0E-110 AB011399.1	ĽΝ	Homo sapiens gene for AF-6, complete cds
12346	L		1.35	1.0E-110		EST_HUMAN	qc31c12.x1 Soares_pregnant_uterus_NbHPU Homo saplens cDNA clone IMAGE:1711222.3'
12429			3.25	1.0E-110		EST_HUMAN	PM3-NN1082-140900-006-f12 NN1082 Homo saplens cDNA
12701	1_		1.45	1.0E-110	1.0E-110 BF508896.1	EST HUMAN	UI-H-BI4-aos-b-05-0-UI.s1 NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:30867843'
170	L		10.84	1.0E-111	1.0E-111 U43701.1	NT	Human ribosomal protein L23a mRNA, complete cds
191	上	25845	1.05	1.0E-111	4758807 NT	LN.	Homo sapiens ras GTPase activating protein-like (NGAP) mRNA
718	1			1.0E-111	11 BF035327.1	EST_HUMAN	601458531F1 NIH_MGC_68 Hamo sapiens cDNA clone IMAGE:3862086 5
726	١	26154	5.13	1.0E-111	8383092 NT	LN.	Homo sapiens cat eye syndrome critical region gene 1 (CECR1), mRNA
906	1_		3.82	1.0E-111	11 M25142.1	IN	Human cardiac alpha-myosh heavy chain (MYH8) gene, exons 32 to 34
1624	1_			1.0E-111	32177	NT	Homo sapiens KIAA0555 gene product (KIAA0555), mRNA
2234	L	L	1.02	1.0E-111	1.0E-111 AF036126.1	NT	Homo sapiens collagen type IX alpha 1 chain (COL9A1) gene, exons 29, 30, 31, and 32
4160	16892		1.08	1.0E-111	7861569 NT	LN T	Homo saplens DKFZP434D156 protein (DKFZP434D156), mKNA
4286	17034		4.38	1.0E-111	1.0E-111 K02268.1	NT	Humen enkephalin B (enkB) gene, exon 4 and 3' flank and complete cds
4691	17426			1.0E-111	TN 8775054	TN	Homo sapiens phosphorylase kinase, alpha 1 (muscle) (PHKA1), mKNA
5644		31249	1.09	1.0E-111	11 BE867909.1	EST_HUMAN	601443690F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3847655 5
	l						qp08g12.x1 NCI_CGAP_Kld5 Homo saplens cDNA clone IMAGE:1917674 3' similar to gc::M28883 KAS- per Atten promitten by La (uli MANN):
5942	18724		1.98	1.0E-111		ES HOMAN	RELATED FROTEIN RALM (TOWAN)
6580	19343	32357	1.16	1.0E-111		EST HUMAN	DKFZp434C1815_r1 434 (symonym: ntess) fromo sapiens culva cione DKFZp434C1919 o
6029	19624	32668	1.06	1.0E-111		EST HUMAN	UI-H-BW0-all-d-03-0-UI.s1 NCI_CGAF_Subs Homo sapiens cDNA ctone IMAGE:2/25020 3
7347	1		2.99	1.0E-111	11 BF366228.1	EST_HUMAN	IL2-NT0101-280700-114-E03 NT0101 Homo sapiens cDNA
7433		33198	0.62	1.0E-1	11 AI761228.1	EST_HUMAN	wi88d01.x/1 NCI_CGAP_Kid12 Homo sepiens cDNA clone IMAGE:2388466 3' similar to gb:J04813 CYTOCHROME P450 IIIA6 (HUMAN);
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Single Exon Plobes Lypresson in State	Top Hit Descriptor	Homo sapiens basic transcription factor 2 p44 (bt/2p44) gene, partial cos, neurona apopusas minimary protein (naip) and survival motor neuron protein (smn) genes, complete cds	G1256410 11-ZINC-FINGER TRANSCRIPTION FACTOR: G1256410 11-ZINC-FINGER TRANSCRIPTION FACTOR:	22/ Egos. 1 NO. CONTROL TRANSCRIPTION FACTOR.; 61266401 1- ZINC-FINGER TRANSCRIPTION FACTOR.; Home sentiens crotein x 000/1 (LOC51185), mRNA	Human beta4-Integrin (ITGB4) gene, excn 13	Homo sapiens nuclear factor of activated T-cells, cytoplasmic 2 (NFATC2), mRNA	Homo sapiens mRNA for FLJ00045 protein, partial cds	Homo sapiens cone sodium-calcium potassium exchanger splice variant (NCKX) mRNA, complete cds	601847132F1 NIH_MGC_55 Homo sapiens cuiva Corio invacativa corio	Human many of integral applied Statement	Human mkny for integral appliance seconds	Homo saprens 110 south 111 CT 11 CT	CASE AND COMP GCR1 Homo saniens cDNA clone IMAGE:825170 3' similar to gb:L09235	aesoguzsi Not_Coart_Coart_Trice. VACUOLAR ATP SYNTHASE CATA TITLE SUBUNIT A, UBIQUITOUS (HUMAN); Li considere DGH1 name retravirus-like element	Homo Sapters No. 11 gard, 1944	Human thrombopoletin receptor (MPL) gene, exons 1,2,3,4,5 and 6		cn07a11.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NH IBC_cn07a11 rangom Long capiens menincipma (disrupted in balanced translocation) 1 (MN1), mRNA	AV70R482 ADC Homo sapiens cDNA clone ADCAOB08 5'	Homo saplens mRNA for neurexin I-alpha protein, complete cds	Homo sapiens acetyl-Coenzyme A carboxylase beta (ACACB), mRNA	Human steroidogenio acute regulatory protein (StAR) gene, exon 5	Human steroidogenic acute regulatory protein (StAR) gene, exon 5	UI-H-BI4-aot-g-04-0-UI.s1 NCI_CGAP_Sub8 Homo saplens cDNA clone IMAGE:3086023 3	UI-H-BI4-act-g-04-0-UI.st NCI_CGAP_Sub8 Homo sapiens cDNA clone IMAGE:3086023 3	Homo sapiens HTRA serine protease (PRSS11) gene, complete cds	ZINC FINGER PROTEIN 135	
le Exon Prop	Top Hit Database Source	Ę	EST_HUMAN	EST HUMAN	Į.		Z	ΤN	EST_HUMAN	۲	Z-	LN	EST HUMAN	EST_HUMAN	TN TOT	ESI HUMAN	ž	EST HUMAN	No. of Property of	ESI TOMAN	124	1 2	2 2	EST LIMAN	EST HIMAN	NT IN	SWISSPROT	
Buis	Top Hit Acession No.	J80017.1	4A278868.1	AA278868.1	1431896	U66533.1	11420310 AK024453.1	AF177987.1		X17033.1	X17033.1	AF091395.1	BF333210.1	AA604160.1			U68159.1	AI75107			AB035356.1	, 0000		-	2 Br-509039.1			
	Most Similar (Top) Hit BLAST E Value	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111	1.0E-111		1.0E-111	1.0E-111			1.0E-11			_		١	١	1		1.0E-112		
	Expression Signal	0.6	0.73	0.73	0.62	3.28	0.79	1.57	8.65	12.9	12.9	2.8	0.49	3.21			4.25											2.2
	ORF SEQ ID NO:	33279	1					34013		34620		١	35066			36038	36902	37376	4 37802									26458
	SEQ ID	20185	1	1	20782		1 1		21399		1	1	l _	1 _			23649	24068	7 24464	l _	17914		L.,		7 13395		1	13800
	Probe SEO ID NO:	7514	7993	7983	8088	8139	8579	8674	8707	0700	8782	8986	71/08	10052	10080	10172	10973	11465	11897	12424	12672	594	596	296	617	617	981	1040

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo seniens K1440/40 motels (K1440440) 1144	Homo sanians KIA 40440 protein (NIA 40440) Fria	IBOTO STATE COAP KIAZ HOME COAP KIAZ	601442674F1 NIH MCC 65 Home capiens con 11 miles and 11 miles and 12 m	Homo sapiens quitamate receptor Involvent Values 4 (CBIXA) — BAIA	Wk45b12.X1 NCI_CGAP_Pr22 Home splens cDNA clone IMAGE:2418335 3' similar to gb:M81660_ms1	MR2-BT0590-090300-143 800 BT0500 (HUMAN);	omo saplens di fometa società società di contra di contr	Homo saplens mRNA for KIA 64444	Homo saplans mRNA for KIAA444 and the saplans mRNA for KIAA4444 and the saplans mRNA for KIAA44444 and the saplans mRNA for KIAA444444 and the saplans mRNA for KIAA4444444444 and the saplans mRNA for KIAA44444444444444444444444444444444444	ASON A Series melanasta SNIL II II protein, partiel cds	Home sanlens NOD1 miles (NOD1)	601564717F1 NIH MCC o Umm	22152648F1 NIH MGC 81 Home collection of the col	6014275551 MIL MCC 44 LL.	6011427555F1 NIH MCC 14 Home sapiens cUNA clone IMAGE:3506508 5	602131405F1 NIH MCC 91 Ums capiens con IIII AC 91 Ums capiens con IIII AC 91 Ums capiens con IIII AC 91 Ums capiens con IIII AC 91 Ums capiens con IIII AC 91 Ums capiens capi	Home equipmental in the september of the	organics source carrier rammy 6 (neurotransmitter transporter, L-proline), member 7 (SLC6A7), mRNA	Homo saplens solute carrier family 6 (neurotransmitter transporter, L-proline). member 7 (SI C6A7). mRNA	601845089F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4070302 5'	AU118051 HEMBA1 Hamo saplens cDNA clone HEMBA1002773 5	601443161F1 NIH MGC 65 Homo sapiens cDNA clone IMAGE:3847285 당	601443151F1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:38472855	730g07.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:3523020 3' similar to	MR3-SNOON9-100-00-10-00-00-00-00-00-00-00-00-00-00	Mo Saplens mRNA for someted modules action to the	601155323F1 NIH MGC 21 Home confere child alone (MA OF access of	qk24c08.y5 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1869902 5' similar to TR:Q64362 Q64362 FUSED TOES;	
gle Exon Prob	Top Hit Database Source			T HUMAN	Г			EST HIMAN				HIMAN		HIMAN	Т	Т	Т	T				7	٦				Т	Т	HUMAN		
ris	Top Hit Acession No.	7662125 NT		AI766925	112 BE866859.1	4504116 NT	112 AI826511.1		24116	112 AB037832.1		Ī	-		Γ	T	Γ		16777		167777	T			12 BE867635.1 E	12 BF111413.1		Γ			
	Most Similar (Top) Hit BLAST E Value	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112	1.0E-112			1.0E-112	1.0E-112	1.0E-112	1.0E-112/				1.0E-112		1.0E-112		1.0E-112	1.05-112 8	A 21 1-2 A	1.0E-112 B	1.0E-112 B	1.0E-112 B	1.0E-112 A	1.0E-112 A	1.0E-112B	1.0E-112 A	
	Expression Signal	4.39	4.39	1.37	1.1	1.15	.0.7	0.74	1.39	4.9	4.9	40.71	1.04	1.43	0.68	0.71	0.71	1.13	1.87	,	1.87	8 6	2/3	80.7	2.09	2.15	2.86	3.37	1.92	1.59	
	ORF SEQ ID NO:	27117	27118	27657	27969		28770	29253	28827	30068	30089	31291	31730	31899	32134	32308	32309	32607	32895	90000	32890	32040	270010	100-0	34695	35649	36611	36796	36981	37053	
	Exen SEQ ID NO:					15842	16116			17437	17437	18378	18766	18932	19138	19304	19304	19575	19921	10021	25420	78702	24547		2164/	22444	23368	23648	23711	23779	
	Probe SEQ ID NO:	1679	1679	2194	2512	3076	3355	3864	4565	4704	4704	5581	5985	6155	6369	6239	6539	6741	7238	7.738	77.30	8083	RREG	3 2	2000	9783	10677	10868	11040	11109	

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77,0,1			-			_	_		_	_										4		*****			7	-uw-	-	****	
	Top Hit Descriptor	qk24c08.y5 NCI_CGAP_Kid3 Homo sapiens cDNA done IMAGE:1869902 5' similar to TR:Q64362 Q64362 FUSED TOES :	PM0-CT0237-141099-001-h02 CT0237 Homo sapiens cDNA	qk24c08.y5 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1869902 5' similar to TR:Q64362 Q64362 PUSED TOES :	qk24c08.y5 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1869902 5' similar to TR:Q64362 Q64362 FUSED TOES;	a095f01 x1 Schiller meningioma Homo sapiens cDNA clone IMAGE:1953625 3'	ac95f01.x1 Schiller meningioma Homo sapiens cDNA clone IMAGE:1953625 3'	Human X-linked phosphoghcerate kinase gene, exon 8	ao95f01.x1 Schiller meningiorna Homo sapiens cDNA clone IMAGE:1953625 3'	Homo sapiens elF4E-transporter mRNA, complete cds	Ui-H-BW1-anl-f-03-0-Ui.s1 NCI_CGAP_Sub7 Homo sapiens cDNA cione IMAGE:3082876 3'	Homo sapiens PLP gene	Homo sapiens mRNA for putative RNA helicase, 3' end	Homo saplens gene for cholecystoldrun type-A receptor, complete cds	Homo sapiens activating transcription factor B (B-ATF), mRNA	Homo saplens activating transcription factor B (B-ATF), mRNA	601469465F1 NIH_MGC_67 Horno saplens cDNA clone IMAGE:3872536 5'	AU127214 NT2RP2 Homo sapiens cDNA clone NT2RP2000807 5	AU140291 PLACE2 Hamo saplens cDNA clone PLACE2000274 5'	Homo sapiens P-glycoprotein (mdr1) mRNA, complete cds	Homo sapiens UDP-N-acety-alpha-D-galactosamine:polypeptide N-acety/galactosaminylransferase 8 (GalNAo-T8) (GALNT8), mRNA	Homo sepiens ATP-binding cassette, sub-family B (MDR/TAP), member 4 (ABCB4), transcript variant B, mRNA	Homo saplens ATP-binding cassette, sub-family B (MDR/TAP), member 4 (ABCB4), transcript variant B, mRNA	Homo sapiens glutamate receptor, lonotropic, N-methyl D-aspartate 2A (GRIN2A) mRNA	Homo saplens glutamate receptor, ionotropic, N-methyl D-aspartate 2A (GRIN2A) mRNA	601152078F1 NIH_MGC_19 Hama saplens cDNA clone IMAGE:3508362 5	601152078F1 NIH_MGC_19 Hamo sapiens cDNA clone IMAGE:3508362 5'	EST371030 MAGE resequences, MAGE Homo saplens cDNA	Homo saplens hypothetical protein FLJ11008 (FLJ11006), mRNA
Top 开	Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	۲	EST_HUMAN	LΝ	EST_HUMAN	LN	NT	LN	N	LN	EST HUMAN	EST_HUMAN	EST_HUMAN	LΝ	ΤN	TN	FZ	N	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN
4	No.	Al792603.1	AW377670.1	AI792603.1	Al792603.1	Al365586.1	A1365586.1	M11965.1	AI365586.1	AF240775.1	BF515218.1	AJ006976.1	AJ223948.1	D85606.1	5453562 NT	5453562 NT	BE780858.1	AU127214.1		AF016535.1	11525737 NT	9961249 NT	D061240	6006002 NT	6006002 NT	BE262161.1	BE262161.1	AW958960.1	8922819 NT
Most Similar	LASTE Value	1.0E-112	1.0E-112	1.0E-112		1.0E-113	1.0E-113	1.0E-113		1.0E-113	Į	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	4 OF 443	1.0E-113	1.0E-113	1.0E-113	1.0E-113		1.0E-113
	Signal	1.59	8.8	1.92	1.92	5.37	5.37	66.7	2.68	1.44	1.02	28.34	1.92	16:0	2.16	2.16	2.97	7.66	4.17	1.47	2.62	0.68	ay C	0.8	0.8	0.78	0.78	0.56	0.48
0	ον σ ον ο:	37054	37085	37734		26152				27382				30300		30336		60608		31589	31725	31806	21807				32980		34629
Exon	SEQ ID NO:	23779	Į	24399	24399	13499	13499	13688	14279	15524	14820	15174	15892	17891		17731	25178	18205	18621	18648	18761	18843	18843	18998	18998	19906	19906	20242	21482
Probe	SEQ ID NO:	11109	11139	11810	11810	725	725	921	1532	1932	2088	2456	3127	4966	5008	2008	5165	5405	5832	5861	5979	6064	A POR	6224	6224	7221	7221	7573	8790

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	801007700E4 hiii 1100	001287708F1 NIH MGC 19 Homo sepiens cDNA clone IMAGE:3827554 5	00128/108F1 NIH_MGC_18 Homo saplens cDNA clone IMAGE:3627864 51	NO I-T I U I 34-28050U-021-d02 F I 0134 Homo sapiens cDNA	rionio sapiens transmembrane protein 2 (TMEM2), mRNA	numen erg protein (ets-related gene) mRNA, complete cds	Homo sapiens KAN binding protein 7 (RANBP7), mRNA	III.HE BNO ALL 40 O III. A WILL SHOWN (KANBP7), MRNA	CITIL DIVINGAN-D-10-0-01.r1 NIH MGC_50 Homo sapiens cDNA clone IMAGE:3077322 6'	602247740F4 NILL MCC 62 11	UI-HF-BNO-gki-b-12-0-UI-I NIH MGC 50 Homo continue CNN 1	hh81e09.y1 NCI_CGAP_GU1 Homo septems cDNA clone IMAGE:2869176 5 similar to TR:060327 060327 KIAA0884 PROTEIN :	hth81a09.y1 NCI_CGAP_GU1 Homo sapiens cDNA done IMAGE:2869176 6' similar to TR:O60327 O60327	NAMOSO FOR I EIN ;	Hours septents grutamente receptor, tonograpic, N-methyl D-aspartate 2A (GRIN2A) mRNA	BO1105820E1 NILL MCC 4.E. L.	ne80b03.r1 NCI_CGAP_GCT Homo sapiens cDNA clone IMAGE:2988366 5' ne80b03.r1 NCI_CGAP_GCT Homo sapiens cDNA clone IMAGE:797089 6' similar to SW:FEN1_HUMAN_ P39728 BI AD ENLICANI IN EASE	nc80b03:r1 NCI_CGE	yd15c01.s1 Soares fetal liver spleen 1NFLS Homo sentlens cDNA clone IMAGE does a chilling in the control of the	gb:A21187 ALPHA-2-MACROGLOBULIN PRECURSOR (HUMAN) contains after reportation along the constitution along the cons	Homo saplens hypothetical protein F_120080 (FI 120080) mBNA	Homo saplens rhabdoid tumor deletion region protein 1 (PTDD1)	Homo sapiens nucleoportn-like protein 1 (NLP 1) mRNA	Homo saplens mRNA for KIAA1276 protein, partial cds	Homo sapiens mRNA for KIAA1278 protein, partial cds	Human gene for catalase (EC 1.11.1.6) exon 2 mapping to chromosome 11 hand 213	601869932F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE-410n214 F	Homo sapiens NOD1 protein (NOD1) gene exons 1.2 and 3	Human interferon-alpha recentor (HaileNalako Box) — hita
lie Exon Pro	Top Hit Database Source	MAN III TOB	TOTAL TOTAL	EST LINAM	NAMOR - PIN	L	-		EST HIMAN	FST HIMAN	FST HIMAN	EST HUMAN	EST_HUMAN		NAMOL		H HI MAAN	I^-	I^{-}	.1	EST_HUMAN							HUMAN		
Julic	Top Hit Acessian No.	BE382842 4	E382842 4	F772987 1	11120267	21535 1	4483007	5453997 INT	W500517 1	691687.1	Γ		1.0E-113 AW630291.1 E		5005	TNICOGNO	292968 1	580720.1	580720.1		0551.1	8923087 NT	7657529 NT	9073				-206374.1 EST	-149773.1 NT	3171.1
	Most Similar (Top) Hit BLAST E Value	1.0E-113	1		1 0F-113	1.0E-113 M	1.0E-113	1.0E-113	1.0E-113 A	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1.0E-113	1 0F-113	1.0E-113	1.0E-113 BE	1.0E-113 AA	1.0E-113 AA		1.0E-114 T7	1.0E-114	1.0E-114	1.0E-114	1.0E-114 AB033102.1	1.0E-114 AB033102.1	1.0E-114 X04086.1	1.0E-114 BF206374.1	1.0E-114 Ai	1.0E-114 J03171.1
	Expression Signal	3.06	3.06	0.93	4	0.45	0.81	0.81	0.61	0.55	0.55	1.83	2.84	2.84	1.39	1.39	2.81	1.32	1.32		6.8	/:/	5.09	4.27	1.28	1.28	2.75	1.02	2.61	0.72
	ORF SEQ ID NO:	34830			35583	35683	35802	35803	36386	36387	36388	37011	37019	37020	31974	31975	37177	37393	37394		26045	20402	26712	27104	25491	25492	28537	78587	28377	29731
	Exon SEQ ID NO:	21682	21682	21968	22381	22481	ı	22598	23160	23161	23161	23737	23746	23746	18998	18998	23890	24082	24082	00,6	2000	9000	14039	14413	1,2071	1/07	13683	7220	16/45	17096
	Probe SEQ ID NO:	8992	8992	9301	9730	9830	9950	9950	10514	10515	10515	11067	11076	11076	11181	11181	11227	11481	11481	930	3 5	2 3		200	7000	700	3128	-	_	4358

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		_					_		_		_								_	- '						<u> </u>			=	<u> </u>	
Top Hit Descriptor	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial ods, alternatively spliced	AV733454 cdA Homo saplens cDNA clone cdABA08 5'	AV733454 cdA Homo sapiens cDNA clone cdABA08 5'	Homo sapiens LIM HOX gene 2 (LHX2) mRNA	Homo sapiens clathrin, heavy polypeptide-like 1 (CLTCL1), mRNA	Homo sapiens TNF-inducible protein CG12-1 (CG12-1), mRNA	Homo saplens hypothetical protein (DJ1042K10.2), mRNA	Homo sapiens hypothetical protein (DJ1042K10.2), mRNA	Homo sapiens HLA-B associated transcript-1 (D6S81E) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	Homo sapiens keratin 16 (KRT18) mRNA	QV4-UM0094-300300-156-b08 UM0094 Homo saplens cDNA	q06f01.x1 NCI_CGAP_GC4 Homo saplens cDNA clone IMAGE:1946809 3' sImiliar to TR:000538 000538	TTF-I INTERACTING PEPTIDE 5;	qt06f01x1 NCI_CGAP_GC4 Homo saplens cDNA clone IMAGE:1946809 3' similer to TR:000638 000638 TTF-I INTERACTING PEPTIDE 5;	Homo saplens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo sepiens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo sapiens ferritin, heavy polypaptide 1 (FTH1) mRNA	Homo sapiens alpha-aminoadipate semialdehyde synthase mRNA, complete cds	Homo sapiens alpha-aminoadipate semialdehyde synthase mRNA, complete cds	Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) genes, complete cds	Homo sapiens KIAA0442 mRNA, partial cds	Homo sapiens testican-1 mRNA, complete cds	QV4-UM0094-300300-156-b08 UM0094 Homo sapiens cDNA	Homo sapiens mRNA for alpha-fubulin 8 (TUBA8 gene)	Homo sapiens mRNA for alpha-tubulin 8 (TUBA8 gene)	Homo sapiens partial TTN gene for titin	Homo sapiens mRNA for KIÁA0350 protein, partial cds	Homo sapiens sir2-like 3 (SIRT3), mRNA	Homo saplens EphA4 (EPHA4) mRNA	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes
Top Hit Database Source	FN	EST HUMAN	EST_HUMAN	F	Z	F	LN L	LN-	LN	⊢Z.	N L	EST_HUMAN		EST_HUMAN	EST HUMAN	1	Z	LN.	F.	Z.	Ļ.	Į.	N _T	EST HUMAN	LZ.	FZ	FZ	L	FZ	LZ	NT
Top Hit Acession No.	14 AF223391.1	14 AV733454.1	14 AV733454.1	4758673 NT	11526317 NT	11418041 NT	11034850 NT	11034850 NT	4758111 NT	4505938 NT	4557887 NT	15 AW804759.1		15 Al339206.1	115 Al339206.1	5174702 NT	5174702 NT	4503794 NT	15 AF229180.1	15 AF229180.1	15 U78027.1	15 AB007902.1	115 AF231124.1	115 AW804759.1	15 AJ245922.1	115 AJ245922.1	15 AJ277892.1	115 AB002348.2	6912659 NT	4758279 NT	115 AL096857.1
Most Similar (Top) Hit BLAST E Value	1.0E-114		1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-115	1.0E-115	1.0E-115	1.0E-115		1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115		1.0E-115					1.0E-115	1.0E-115	1.0E-115				
Expression Signal	1.71	9	9	1.7	1.32	3.42	4.93	4.93	2.89	2.03	2.33	2.23		0.99	66.0	1.36	1.36	40.4	1.26	1.26	101	0.98	2.13	1.39	6.22	6.22		3.67		3.6	2.64
ORF SEQ ID NO:	36715				37759					25585		25733		25939	076940	26201	28202		L	26986			27758	l	28518			29397	L	L	Ц
Exon SEQ ID NO:	23472	L	L	24386	L	L		24902	12849	12942	12948	L	L	13307	13307	L	L	L			<u>l</u>		1_	L	15878	L	L	L	L	L	Ш
Probe SEQ ID NO:	10789	11145	11145	11796	11834	12334	12577	12577	21	127	131	286		523	523	769	769	Ě	1552	1552	1833	2078	2298	2855	3113	3113	3465	4021	4369	4403	4628

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Top Hit Descriptor	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes	Homo saplens chromosome 21 segment HS21C068	Homo sapiens chromosome 21 segment HS21C068	EST382416 MAGE resequences, MAGK Homo sepiens cDNA	602119346F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4276738 5'	Homo sapiens similar to ER to nucleus signalling 1 (H. sepiens) (LOC63433), mRNA	Homo sapiens similar to ER to nucleus signalling 1 (H. sapiens) (LOC63433), mRNA	au64g01.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2519568 3' similar to gb1.07897 DYNAMIN-1 (HUMAN);	au64g01.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2519568 3' similar to gb:L07807 DYNAMIN-1 (HUMAN);	Homo saplens sperm surface protein (HSS), mRNA	Homo saplens sperm surface protein (HSS), mRNA	Homo sapiens similar to ribosomal protein S26 (H. sapiens) (LOC63436), mRNA	Hamo saplens KIAA0054 gene product; Helicase (KIAA0054), mRNA	Homo sapiens KIAA0054 gene product; Helicase (KIAA0054), mRNA	yd86b08.r1 Sogres fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:115095 6' similar to SP:DPOG_YEAST P15801 DNA POLYMERASE GAMMA;	oz31a08.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA-clone IMAGE:1676914 3'	oz31a06.x1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:1676914 3'	Homo sapiens mRNA for KIAA0995 protein, partial cds	RC6-ET0081-130700-011-G01 ET0081 Homo sepiens cDNA	RC6-ET0081-130700-011-G01 ET0081 Homo sapiens cDNA	Homo sapiens eukaryotic translation trittation factor 4B (EIF4B), mRNA	601816352F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4050108 5'	Human mRNA for KIAA0338 gene, partial cds	Human mRNA for KIAA0338 gene, partial cds	qg99e09.xf Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1843336 3'	qg99e09.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1843336 3'	#12a07.x1 NCI_CGAP_CLL1 Home sapiens cDNA clone IMAGE:2118036 3' similar to TR:O16129 O16129 PHENYLALANYL TRNA SYNTHETASE;	xx32f08.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2839239 3' similar to SW:CAYP_CANFA P10463 CALCYPHOSINE;	Homo sapiens UDP-ducose:glycoprotein glucosyltransferase 1 (HUGT1), mRNA	
Top Hit Database Source	N	Ŋ	님	EST_HUMAN	EST_HUMAN	N	N F	EST_HUMAN	EST HUMAN	N	NT.	L	F	Ľ.	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	П	EST_HUMAN	N	EST_HUMAN			EST_HUMAN	П	EST_HUMAN	EST HUMAN	NT.	
Top Hit Acessian No.	5 AL096857.1	5 AL163268.2	5 AL163268.2	15 AW970335.1	Γ	15 11425128 NT	11425128 NT	5 A1928799.1	5 A1928799.1	11426786 NT	11426786 NT	11426038 NT	7661883 NT	7661883 NT	5 T86774.1	5 AI076598.1		5 AB023212.1		5 BE830187.1	11434772 NT	5 BF382029.1			5 AI221878.1	5 AI221878.1	5 AI524687.1		15 8910279 NT	
Most Similar (Top) Hit BLAST E Value	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	
Expression Signal	2.64	3.51	3.51	1.62	0.78	1.96	1.96	1.34	1.34	76.0	76.0	9.84	2.04	2.04	0.83	1.54	1.54	8.22	13.71	13.71	2.15	0.6	2.25	2.25	1	1	0.82	7.62	1.33	
ORF SEQ ID NO:	28897	30201	30202	30698	30802	31143	31144	31313	31314	31916	31917	32061	32198	32199	32521	32935	32836	33068	33885	33886	34548	35522	35743	35744	36284	36285	36293	36549	36797	
SEQ ID NO:	17363	17578	17578	18069		18253	18253	18400	18400	18945	18945	19075	19202	19202	19497	19864	19864	19991	20754	20754	21404	22327	22549	22549	23064	23064	23072	23310	23549	
Probe SEQ ID NO:	4628	4848	4848	5263	5338	5454	5454	5604	5604	6168	6168	6302	6434	6434	6835	7178	7178	7308	8060	8060	8712	9675	6686	9899	10418	10418	10426	10617	10869	

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		38378 088378	18378 088378				(GSTT1)									2) gene and	_		ř				element	of solimitor		Ī		Î
Single Exon Probes Expressed in Brain	Top Hit Descriptor	hq54c10.x1 NCI_CGAP_Pan3 Homo sapiens cDNA clone IMAGE:3123186 3' similar to TR:086378 O86378 PRP4 PROTEIN KINASE HOMOLOG	hq54c10.x1 NCI_CGAP_Pan3 Home saplens cDNA clone IMAGE:3123186 3' similar to TR:088378 088378	Home saniers coloins shows 1.16	60111744FI NIH MCC 18 Home College Copendent, alpha 1E subunit (CACNA1E) mRNA	QV3-070065-280300-137-h12 OT0065 Home septems cDNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	601121347F1 NIH MCC 20 Hamman FRIA	Homo sapiens synapticiant 1 / SWI 11 - DNA	Homo sapiens synaptianing (SYN II) = DNA	Homo sablens pericentrin (PCNT) BNA	Homo sapiens nericentry (PCNT) BNA	Human apolinomiain 8 100 (mms)	Human apolipopmiein B-100 (apoe) yene, exons 17 and 18	Homo sablens moteln phospheters, Efficial and the firm	Human olfactory receptor off17-201-1 (OR17-201-1) gene, olfactory receptor olf17-32 (OR17-32) gene and	Hams conjust the DNA (17,000 of 17,000) pseudogene, complete cds	POTESTATES MINING TO NIGAU/BU protein, partial cds	Homo saplens Difference and from the suprementation of the limited	Homo sepiens Diocaca ge syndrome critical region, centromeric end	Homo saniene sodium shock to the characteristics of the contractions and the characteristics of the characteristic	PW-BT135-070499-016 BT135 Home sealers of NPT 3) mRNA	qn19d04.x1 NC _CGAP_Lu5 Homo saplens cDNA clone IMAGE:1898895 3' cimilar to contains element MER25 repetitive element	222407.r1 Sogres_senescent_fibroblasts_NbHSF Homo sapiens cDNA clone IMAGF:322246 From Inch	SW::MDHM_MOUSE P08249 MALATE DEHYDROGENASE, MITOCHONDRIAL PRECURSOR	Homo saplens mRNA for KIAA1636 protein, partial cds	Homo saplens mRNA for KIAA1636 protein, partial cds	801302281F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3638764 F
gle Exon Pro	Top Hit Database Source	EST_HUMAN	FST HIMAN		EST HUMAN	EST HUMAN	L	EST HUMAN	LZ	F								T HIMAN	Т			T HUMAN			HOMAN			EST HUMAN 6
Sin	Top Hit Acession No.	15 BE045890.1	115 BE045890 1	4502528 NT	115 BE255549.1		15 AF240786.1		07334	4507334 NT	5174478 NT	5174478 NT	16 M19824.1		5453941	16 U78308 1	-				5031954	16 A1907096.1	I6 AI302062.1 E		10 W 42822.1		T	o BE408097.1 E
	Most Similar (Top) Hit BLAST E Value	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115 /	1.0E-115	1.0E-116	1.0E-116	1.0E-118	1.0E-116	1.0E-116	1.0E-116 N	1.0E-116 N	1.0E-116	1.0E-116 U	1.0E-118 A	1.0E-118 B	1.0E-116 L77570.1	1.0E-116 L.	1.0E-116	1.0E-116 AI	1.0E-116 AI	L	1.0E-110 W	1.0E-116 AE	1.0E-110 AE	1.0E-116 BE
	Expression Signal	1.54	1.54	2.27	2.53	1.63	2.16	1.98	2.21	1.78	2.55	2.55	2.6	2.6	1.95	1.36	2.84	1.53	4.87	4.87	2.43	1.57	0.87	;	7	0. 0	0.78	0.70
	ORF SEQ ID NO:	37174	37175	37281	37699	37767		25969	26216		27447	27448	27552	27553	27767		27915	28183	28582	28583	29713	30156	30627	26	34882	31863	31834	2
	Exon SEQ ID NO:	23888	23888	23981	24366	24426	24481	13341	13555	13609	14726	14726	15585	15585	15030	15063	15176	15533	15934	16934	17084	17534	18005	18674	18895	18895	18961	3 5
	Probe SEQ ID NO:	11225	11225	11374	11775	11842	11920	629	88	839	<u>8</u>	1990	88	2080 2080	2305	2340	2458	2738	3171	3171	4345	8 803	5197	5880	6117	6117	9184	200

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Top Hit Descriptor	C02944 Human heart cDNA (YNakamura) Homo saplens cDNA clone 3NHC0567	AV716314 DCB Homo saplens cDNA clone DCBBCG06 5	EST62685 Jurkat T-cells V Homo saplans cDNA 5' end similar to similar to keratin 2	EST62685 Jurkat T-cells V Homo sapiens cDNA 5' end similar to similar to keratin 2	CM-BT043-090299-075 BT043 Homo sapiens cDNA	601338268F1 NIH_MGC_53 Homo sapiens cDNA clone IMAGE:3680680 5'	qh09c05.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA done IMAGE:1844168 3' similar to gb:X53741 ma1 FIBULIN-1, ISOFORM A PRECURSOR (HUMAN);	Homo saplens laminin, alpha 2 (merosin, congenital muscular dystrophy) (LAMA2), mRNA	Homo sapiens partial mRNA for xylosyltransferase I (XT-I gene)	Homo sapiens partial mRNA for xylosyltransferase I (XT-I gene)	QV4-HT0401-281299-063-c09 HT0401 Homo sepiens cDNA	CM2-CT0482-300800-349-e06 CT0482 Homo sapiens cDNA	qq41e04.x1 Soares_NhHMPu_S1 Homo sepiens cDNA clone IMAGE:1835102 3' similær to WP:B0495.7 CE01765:	DKFZp762L1110_r1 762 (synonym: hmel2) Homo saplens cDNA clone DKFZp762L1110 5'	Homo sapiens acety-Coenzyme A carboxylase alpha (ACACA), mRNA	Mus musculus fragile-X-related protein 1 (Fx1h) gene, exons 13a through 15	Homo sapiens lymphocyte activation-associated protein mRNA, complete cds	Human apolipoprotein B-100 (apoB) gene, exon 10	EST369769 MAGE resequences, MAGE Homo saplens cDNA	op32c11.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1578548 3'	EST188414 HCC cell line (matastasis to liver in mouse) II Homo sapiens cDNA 6' end similar to ribosomal protein L29	Homo sapiens collagen, type IV, alpha 5 (Alport syndrome) (COL445), mRNA	DKFZp434C1120_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C1120 5'	H. sapiens mRNA for TPCR16 protein	H.sapiens mRNA for TPCR16 protein	Homo sapiens Scar2 (SCAR2) gene, partial cds	Homo sapiens Scar2 (SCAR2) gene, partial cds	Homo sapiens mRNA for KIAA0866 protein, complete cds	Homo sapiens atrophin-1 interacting protein 1; activin receptor interacting protein 1 (KIAA0705), mRNA	601562857F1 NIH_MGC_20 Hamo sapiens cDNA clane IMAGE:3832214 5'
Top Hit Database Source	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	LN	⊢Z	N	EST_HUMAN	EST_HUMAN	EST HIMAN	EST HUMAN	LZ	L	L L	LN	EST_HUMAN .	EST_HUMAN	EST HIMAN	LN	EST_HUMAN	LN	N	LN	۲N	IN	NT	EST_HUMAN
Top Hit Acession No.	16 C02944.1	16 AV716314.1	16 AA354258.1	16 AA354258.1	16 Al904151.1	16 BE565507.1	16 AI218352.1	11418646 NT	16 AJ277441.1	16 AJ277441.1	1.0E-116 BE158913.1	16 BF335849.1	16 A1367140 1	16 AL134889.1	4826836 NT	17 AF124393.1	1.0E-117 AF123320.1	17 M19816.1	17 AW957699.1	1.0E-117 AA978114.1	17 44318723 1	8659564 NT	17 AL042120.1			1.0E-117 AF134304.2	17 AF134304.2	17 AB020673.1	6912461 NT	17 BE730508.1
Most Similar (Top) Hit BLAST E Value	1.0E-116	1.0E-118	1.0E-118	1.0E-118	1.0E-116	1.0E-116	1.0E-116		1.0E-116	1.0E-116	1.0E-116	1.0E-116	1 0F-118		1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1 0F-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117 X89670.1	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117
Expression Signal	1.19	5.74	1.37	1.37	0.98	1.66	1.61	1.52	0.74	0.74	0.78	2.4	285	1.86	1.67	96.0	1.02	1.51	1.54	1.64	2.1	2.03	2.95	1.27	1.27	10.03	10.03	3.57	0.73	3.01
ORF SEQ ID NO:	32572		34101	34102		34674	34837				38104		32025		25959											30119	30120	30233	30471	
Exen SEQ ID NO:	19544	19790	20961	20961	21071	1	21687	1	1	22818	22894		23750	25267	13328	15559	14489	14582	14936	16024	16720	17049	17273	17408	17408	17491	17491	17614	17854	ΙI
Probe SEQ ID NO:	6844	7102	8267	8267	8278	8836	8997	. 8573	10170	10170	10248	10586	11080	12625	545	1055	1747	1823	2208	3262	3974	4310	4538	4674	4674	4759	4759	4887	5136	5284

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Top Hit Descriptor	EST26111 Cerebellum II Homo sapiens cDNA 5' end similar to similar to zinc finger domain	Homo sapiens nuclear hormone receptor (shp) gene, 3' end of cds	Homo sapiens nuclear hormone receptor (shp) gene, 3' end of cds	AV717788 DCB Homo seplens cDNA clone DCBBAE01 6	AV717788 DCB Homo sapiens cDNA clone DCBBAE01 5'	wp86b07.x1 NCI_CGAP_Brn25 Hamo sepiens cDNA done IMAGE:2468629 3 similar to 1 R:O/3065 075065 KIAA0477 PROTEIN.;	Homo sapiens neural cell adhesion molecule 1 (NCAM1), mRNA	Homo sapiens neural cell adhesion molecule 1 (NCAM1), mRNA	CM-BT043-090299-075 BT043 Homo saplens cDNA	CM-BT043-090299-075 BT043 Homo saplens cDNA	Human gene for very low density lipopratein receptor, exch 11	601569317F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3843748 5	Homo sapiens gamma-aminobutyric acid type B receptor 2 (GABABK2) mKNA, complete cos	Homo sapiens Drosophila Kelch like protein (DKELCHL), mRNA	Human mRNA for KIAA0191 gene, partial cds	Homo sapiens protein (peptidy-proty dis/trans isomerase) NIMA-interacting 1 (Pin 1), minuk	Homo saplens protein (peptidyl-prolyl cis/trans isomerase) NiMA-interacting 1 (PIN1), mKNA	Homo sapiens mRNA for MEGF8, partial cds	Homo saplens mRNA for MEGF8, partial cds	601186203F1 NIH_MGC_8 Hamo sepiens cDNA dane IMAGE:3344289 3	Homo saplens ATP-binding cassette, sub-tamily A (ABC.1), member 3 (ABCA3), mixing	Homo sapiens ATP-binding cassette, sub-tamily A (ABC1), member 3 (ABCA5), mrNA	Homo sapiens HSPC151 mrkNA, complete cas	DKFZp4341056 71 434 (synonym: ntess) Homo sapiens cunA cione DNFZp4341050 0	Homo capiens hypothetical protein (DJ320ET 19.01.1), infinity and infi	Hamo sapiens sine oculis nonecoox (Urasophila) nonecoox (Urasophil	601281947F1 NIH MGC_44 Homo sapiens cDNA clone IMAGE:3504018 5	601281947F1 NIH_MGC_44 Homo sapiens cDNA clone limAGE:3504018 3	60128194771 NIH MGC 44 Home saplens cUNA clone INACE: 3004019 o	EST363799 MAGE resequences, MAGB Homo sapiens cDNA	Human breakpoint cluster region (BCR) gene, complete cds	Human breakpoint cluster region (BCR) gene, complete cas	Homo Bapiens Print Excit
Top Hit Database Source	EST_HUMAN	LN TN	N	EST_HUMAN	EST_HUMAN	EST HUMAN	N L	Į.	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	TN	NT	NT	NT	NT	LN	NT	EST_HUMAN	N	NT	NT	EST HUMAN	Į.	N	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	۲	LV.	Ł
Top Hit Acession No.	17 AA323348.1	17 L76571.1	17 L76571.1	17 AV717788.1	17 AV717788.1	17 AI950145.1	10834989 NT	10834989 NT	17 AI904151.1	17 A1904151.1	17 D16524.1	17 BE733922.1	17 AF099033.1	11420222 NT	17 D83776.1	11424835 NT	11424835 NT	17 AB011541.1	17 AB011541.1	17 BE269856.1	4501848 NT	4501848 NT	18 AF161500.1	AL04585		5174680 NT	18 BE389705.1	18 BE389705.1	18 BE389705.1	18 AW951729.1	18 U07000.1	118 U07000.1	18 713932.1
Most Similar (Top) Hit BLAST E Value	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1 0F-117		1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-117	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118		1.0E-118
Expression Signal	0.86	5.01	5.01	1.75	1.75	àc c	220	229	0.56	0.56	2.25	2.07	2.9	1.11	1.77	2.68	2.68			14.73	2.02	2.02	5.98	2.13	5.67	96.0	2.04	2.04	2.04	1.68	2.61		4.64
ORF SEQ ID NO:	30574	l								L			L					l	37197		37501	37502	25530		25922	26328		27694			28189		
Exan SEQ ID NO:	17937	1		1	ſ	1	L	1	1	L	L	L			<u> </u>	Ł.	L	Ł	L	L	1	L	12896	L	13288	15555	L	L	L		1	1	1 1
Probe SEQ ID NO:	6860	7350	7350	7448	7446	9	0/0/0	8210	8310	8310	9189	9896	9846	10469	10765	10964	10964	11243	11243	11369	11687	11587	8	9	52	8	2227	2227	2227	2328	2744	2744	3102

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					A I	Siligle Exoli Floor	
Probe SEQ ID	SEO ID	ORF SEQ ID NO:	Expression	Most Similar (Top) Hit BLAST E	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
Š	<u>.</u>			value		Т	A SASE TA NOT COAD KINS Home seniens cDNA clone IMAGE:19167693'
3190	15953	28605	4.67	1.0E-118	Al347694.1	Т	appullobix I NOT COSAP Kid5 Home saplens cDNA clone IMAGE:1916769 3'
3190			4.67		AI347694.1	HOMAN	gpo Hossia in National protein, complete cds
4067	16811	29439	4.77	1.0E-118	D23660.1	Z	Hamp sablens KIAA0478 gene product (KIAA0478), mRNA
4859	1_	30028	6.0		11425/93	2	industrial calcium channel damma 4 subunit (CACNG4) gene, exon 3
5337	L	30800	1.87	1.0E-118	AF142624.1	Z	Trail organization channel damma 4 subunit (CACNG4) gene, exch 3
5337			1.87	1.0E-118	AF1426	LN.	Home salens reein (RELN), mRNA
5543	_	31247	0.94			Z	Homo saplens reelin (RELN), mRNA
6543	18340	31248			11422054	2	Human custic flurosis transmembrane conductance regulator (CFTR) gene, exon 4
5684	L	31395	1.24		M55109.1	Z	Hame agreem T-box 4 (TBX4), mRNA
5772	١.	L	0.83			Z	Harris September 7-hove 4 (TBX4) mRNA
6777	1	L	0.83			L	Hours experient recenter potential channel 5 (TRPC5), mRNA
5853	1_		1.49	1.0E-118	_	L	Home sapiraris usus mensforming growth factor beta binding protein 2 (LTBP2) mRNA
2000	1					N	Homo sapiens laterit danscrining a committee beta binding protein 2 (LTBP2) mRNA
0032 0502	1.				4557732 NT	Ę	Homo sapients latent transformming & Thomas Sapiens cDNA clone DKFZp43400127 5
	l			1.0E-118	3 AL043761.1	EST HUMAN	UKFZP84H0127_1 - 1-4-(Spring) Here (Proposed Pro
686	L	1			1.0E-118 AL043761.1	EST HUMAN	DKFZp434C012/_T1 454 (synonym: mess) riching (C2ORF3), mRNA
	L			L	3 11431050 NT		Homo saplens curomoscana z open recently preme INACEF 4302749 5
7500	\perp	l			BF6852	EST_HUMAN	602141620F1 NIH MGC 49 Home suppers of the man from the Figure 100 ft.
7,33	20397	33512			1,0E-118 BE781223.1	EST HUMAN	601469169F1 NIH MGC_67 Home sapiens clink civile with Clinks
<u> </u>	_				8 BE062855.1	EST_HUMAN	QVQ-BIQ2SJ-U9UZUJ-91-110- DI VZCST I TOTIO SERIENTE CINA
8282				1 0E-11	8 BE062855.1	EST HUMAN	QVQ-B10283-090200-091-700- B10280-1040-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0
8282	_1				8 AA443024.1	EST_HUMAN	2x98d07.r1 Soares_NhHMPu_S1 Home sapiens CDNA clotte in Active 1/1789 5
8288	_1			L	1 0F-118 AA443024.1	EST_HUMAN	268d07.1 Scares NhHMPu S1 Home saperis Contractions
8288	L	1		1 OF-11	8 AB002381.1	LN	Human mRNA for KIAA0383 gene, per usi
8573	- !			1 10	B AB002381.1	FN	Human mRNA for KIAA0383 gene, partial cos
85.				1-10-1	TN CCASSON	LN.	Homo sapiens latent transforming growth factor beta binding proper 2 (1 or 2) in the contract of the contract
8621	21 21313			1.0E-17		T.V.	Homo saplens latent transforming growth factor beta binding protein 2 (L i BF2) mixina
8621	21 21313	13 34456			8	FOT LIMAN	R01144863F2 NIH MGC 19 Homo saplens cDNA clone IMAGE:3160502 5
8933	33 21624	24 34767		1.0E-17	1.0E-118 BE263134.1	NAME IN THE	DKFZp588K1824 rt 588 (synonym: hute1) Homo saplens cDNA clone DKFZp586K1824
8064	L	55 34806	6 0.55	1.0E-1	B AL04847	ESI DOMEN	Limen contains hymothetical protein (DJ328E19.C1.1), mRNA
3					18 7657016 N1	NAME TO FOUR	HRITO CAPACION 1 547 (synonym: hfbr1) Homo sapiens cDNA clone DKFZp5470017 5
8	<u>l</u>	36 35731	1 0.98	.	1.0E-118 AL138321.1	NONDE I CE	7417e/09 x1 NCI CGAP Brn23 Homo sapiens cDNA clone IMAGE:3564785 3' similar to SW:ZP3A_HUMAN
10024	1	36134	1.88	1.0E-1	18 BF195407.1	EST_HUMAN	\neg
104	١	1					

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	xe46e10.x1 NCI_CGAP_Kid11 Homo septiens oDNA clone IMAGE:2772666 3' similar to	SW. BOLDG THOMAN OVESSE GAMMA-BUTYROBETAINE, 2-OXOGLUTARATE DIOXYGENASE;	Grand Strate Control Mac Control Strate Strate Control Strate Control Mac Cont	oce 11 tezen 1 km _ woo to riomo septens cDNA clone IMAGE:4302798 6* Homo septens protein with polyglutamine repeat, calcium (ca2+) homeostasis endoplasmic reticulum protein (ERPRO 7243-21) mRNA	EST186814 HCC cell line (matastasis to liver in mouse) Il Homo sepiens cDNA & end similar to dynein, light	Shadogada Nilo Moo as II	ISO1489614E1 NIT WAS 19 TOTAL SEPTENS CON CIONE IMAGE:3901563 5	OVO-UMM091-1200-1-1 INIT_INITEC_/ O Home sapiens clone IMAGE:3901563 5	OVO-UMORO1-120000-38E-1-3 III Appear 1-2	Homo espiens children chame of C4 (AICA) - BM14	Homo seniens CGL-105 projets (1 OC# 204) mrnvA, complete cas	Homo sariens mRN & for KTA Angas	Home saplens handhelical braken El 140052 / El 140052 / El 140052	on10b05.s1 NCI_CGAP_Lu6 Home sapiens cDNA clone IMAGE:1566241 3' sImilar to WP:E04F6.2	Use (214)	From Septems glutamete receptor, longropic, kainate 1 (GRIK1) mRNA	Himan native from A NEW Y Hours Spiens CUNA clone NT2RP4001991 6	RC1-NN0073-250800-018-08 NN0073 House cas	AV693731 GKC Home sablens cDNA clans CKCDDB02 #1	DKFZp762M0710 r1 782 (swonym: hme/2) Home conical - BN/A - 1	DKFZp762M0710 r1 762 (synonym: hmel2) Homo septems cDNA clone DKFZp762M0710 6	4b77c09.x1 Soares_fetal_heart_NbHHJ9W Homo saplens cDNA clone IMAGE:1709128 3' elmilar to	Homo septems matrix matally and the CYTOSKELETAL 10;	Homo saplens matrix matellacentalises 28 (MMP28) mRNA, complete cds	m23ff0 vf Scenes NET 7 CBC 64 H	Human creeting protections.	EST386298 MAGE resenience: MAGM Home control	801692005F1 NIH MGC 7 Homo sanlens o'DNA chara IVA CE-2046084 ET	601280564F1 NIH_MGC_39 Home sapiens cDNA clone IMAGE:3822526 5
gle Exon Pro	Top Hit Database Source	FOT LI MAN	TO TOWN	EST HIMAN	TN	MANNI TO THE	EST HIMAN	EST HIMAN	EST HUMAN	EST HUMAN	Z		LN L	7		NEW POLICE	T HIMANI	Т	HOMAN	Г	Т	Т		Т		T HUMAN	Т	T HUMAN		П
uis	Top Hit Acession No.	-118 AW277289 1	.118 AW296351 1	118 BF685214.1	11055968 NT	118 AA315007 1	118 BE908676.1			Γ		7705607 NT	19 AB023147.1	22205	19 44916780 4	4116	19 AU133389 1		-	19 AV693731.1 E	19 AL134903.1 E	19 AL 134903.1 E	19 A1150703.1		Τ	T		3.1		19 BE615150.1 E
	Most Similar (Top) Hit BLAST E Value	1.0E-118	1.0E-118	1.0E-118	1.0E-118	1.0E-118		1.0E-118	1.0E-118	1.0E-118	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0F-119	1.0E-119	1.0E-119	1.0E-119 M89914.1	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119 A		1.0E-119 AI476732.1	1.0E-119 X06292.1	1.0E-119 A	1.0E-119 B	1.0E-119B
	Expression Signal	0.46	0.65	1.61	1.8	10.23	1.68	1.68	1.61	1.61	0.97	1.61	5.97	1.57	8.0	1.42	2.5	21.82	3.11	1.83	0.63	0.63	7.67	0.92	0.92	0.85	2.62	4.69	1.5	1.19
	ORF SEQ ID NO:	36227		37,155	37186	37201	37457	37458	37461	37462	26173	26433	27374	28506		29325	30688	30701	30706	30830	31201	31202	31778	31940	31941	31987	32135	32148	33070	34390
	Exan SEQ ID NO:	23012	23077	23869	23899	23908	24147					-	- 1	15864	15996	16684				- 1	18301	18301	18816	18967	18967	19013	19139	19149	19993	21252
	Probe SEQ ID NO:	10365	10431	11206	11236	11246	11548	11548	11551	11551	741	1014	1928	3089	3234	3934	6253	5266	5270	5347	2203	9203	8038	6190	6190	6239	6370	6380	7310	8260

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	Γ		Π	Γ	Г		Γ	Г	Г	Г	Г		Г	Г	Γ		Γ	٢					Г		П	Γ	Г	<u> </u>		Г			П
Top Hit Descriptor	Homo sapiens KIAA0477 gene product (KIAA0477), mRNA	qf43a11x1 Sogres_testis_NHT Homo saplens cDNA clone IMAGE:1752764 3' similar to TR:Q13458 Q13458 GUANINE NUCLEOTIDE EXCHANGE FACTOR PROTEIN TRIO.;	aa32005.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:814977 5'	Homo sapiens partial IL-12RB1 gene for IL-12 receptor beta1 chain, exons 16-17	Homo sapiens hypothetical protein FLJ10208 (FLJ10208), mRNA	Homo sapiens hypothetical protein FLJ10206 (FLJ10206), mRNA	Homo sapiens Sod mRNA for stearcyl-CoA desaturase, complete cds	Homo sapiens partial IL-12RB1 gene for IL-12 receptor beta1 chain, exons 16-17	Homo sapiens partial IL-12RB1 gene for IL-12 receptor beta1 chain, exons 16-17	602186072F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4310633 5'	RC3-CT0212-240999-011-f03 CT0212 Homo sapiens cDNA	Homo sapiens synaptojanin 1 (SYNJ1), mRNA	Homo saplens intersectin 2 (SH3D1B) mRNA, complete cds	Homo saplens intersectin 2 (SH3D1B) mRNA, complete cds	yy40g12.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:273766 5'	Homo sapiens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sapiens disintegrin and metalloprotease domain 10 (ADAM10) mRNA	Homo saplens gene for AF-6, complete cds	Homo sapiens gene for AF-6, complete cds	Homo sapiens aquaporin 4 (AQP4), splice variant b, mRNA	Homo sapiens synaptojanin 1 (SYNJ1), mRNA	Homo sapiens cAMP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	Homo sapiens cAMP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	Homo saplens stanniocalcin (STC) gene, partial cds	Homo sapiens stanniocalcin (STC) gene, partial cds	Homo saplens glutamate receptor, bnotroplo, kalnate 1 (GRIK1) mRNA	qd61f03.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1733981 3'	602183994F1 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300174 5'	602183994F1 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300174 5'	Human P-glycoprotein (MDR1) gene, exons 6 and 7	Human P-glycoprotein (MDR1) gene, exons 6 and 7	Human TBXAS1 gene for thromboxane synthase, exon 7	Human gene for neurofilament subunit M (NF-M)
Top Hit Database Source	TN	EST HUMAN	EST_HUMAN	TN	NT	NT	IN	NT	NT	EST_HUMAN	EST_HUMAN	TN	N.	L	EST_HUMAN	N	TN	NT	NT	NT	NT	TN	LΝ	LN	LN	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	NT	LN
Top Hit Acession No.	11036643 NT	9 Al149796.1	9 AA465124.1	9 AJ297701.1	11425837 NT	11425837 NT	1.0E-119 AB032261.1	1.0E-119 AJ297701.1	1.0E-119 AJ297701.1	9 BF569571.1	9 AW847519.1	4507334	1.0E-120 AF248540.1			1.0E-120 AF167706.1	4557250 NT	0 AB011399.1	AB011399.1	4755124 NT	4507334 NT	AF056490.1	AF056490.1	AF098463.1	AF098463.1	4504116 NT	A/190903.1		:1	M29428.1	M29428.1		
Most Similar (Top) Hit BLAST E Value	1.0E-119	1.0E-119		1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-119	1.0E-120	1.0E-120	1.0E-120 A	1.0E-120 N44873.1	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120 A	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120 A			1.0E-120 N	1.0E-120 N	1.0E-120	1.0E-120 Y00067.1
Expression Signal	1.15	0.55	3.35	1.29	0.71	0.71	4.16	2.38	2.38	6.31	2.16	1.43	2.49	2.49	2.31	3.08	1.21	0.92	0.92	5.24	1.59	1.95	1.95	2.22	2.22	1.36	6.0	16.61	16.61	0.57	0.57	1.77	6.22
ORF SEQ ID NO:	35663	35867	36008	36263	36317	36318	06898	36913	36914			25741	26439	26440	26832	27033	27250	27565	27666	27985	25741	29692	29693	29983	29984	30401	30468	31357	31358	32110	Ш		33612
Exan SEQ ID NO:	22458	22654	L	23047	23089	23089	23163	23660	23660	23826	25326	13100	13778	13778	14152	14343	14539	14831	14831		13100	17064	17064	17349	17349	17784	17851			l	11		20490
Probe SEQ ID NO:	9807	10006	10146	10401	10443	10443	10517	10985	10985	11159	12198	294	1018	1018	1405	1597	1799	2100	.2100	2531	3302	4325	4325	4614	4614	2002	5133	5649	5649	6350	6350	7471	7795

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	Top Hit Descriptor	- 57-04 -4 NCI CraP Pan1 Homo sapiens cDNA clone IMAGE:2005417 3'	distriction of the few	Findens of NCI CGAP Lu24 Home sapiens cDNA clone IMAGE:3166119 3	A01140485F1 NIH MGC 9 Homo sapiens cDNA clone IMAGE:3049820 5	Himan alucose transporter (GLUT4) gene, complete cds	Homo saplens Xg pseudoautosomal region; segment 2/2	RC3-NN0068-270400-011-f02 NN0068 Homo sapiens cDNA	RC3-NN0066-270400-011-f02 NN0066 Homo sapiens cDNA	Homo saciens gamma-aminobutyric acid (GABA) A receptor, alpha < (CADINAE), III. C.	Homo sapiens DNA for prostacyclin synthase, exon 8	Homo saplens DNA for prostacyclin synthase, exon 8	in Scots vi Human Pancreatic Islets Homo saplens cDNA 5' similar to 1 K: 0/345/ 0/345/ 0/1000	PHOSPHOLIPASE A2-GAMMA. : increme of Human Pancreatic Islets Homo sapiens cDNA 5' similar to TR:075457 075457 CYTOSOLIC	RECOGNIC PROTECTION OF THE PRO	Homo seriens UDP-glucuronosyltransferase 2B4 precursor (UGT2B4) mRNA, UGT2B4*E458 allele,	complete cds	Homo sapiens critolide intracollular and a sapiens con IMAGE:248448 3'	AU119320 HEMBA1 Homo saplens cDNA clone HEMBA1005536 5	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1), mrtiva	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens T-cell lymphoma invasion and metastasis 1 (I LAWI I), III N.S.	Homo saplens intersectin short isoform (115N) mKNA, comprete cas	(Il segnous V kappa II)	Human kappa-immunoglobulin germline pseudogene (Crizz) sarang saperalmunoglobulin germline protein SS2 precursor, mRNA, complete cds Homo sapiens cysteine-rich repeat-comfaining protein SS2 precursor, mRNA, complete cds	Homo sepiens collagen, type XII, alpha 1 (COL1241), mKNA	Homo saplens collagen, type XII, alpha 1 (COL1241), mRNA	601497032F1 NIH_MGC_70 Homo sapiens cDNA clone IMAGE: 3038330 3	601896173F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE: 4126234 5	601896173F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4125234 3	
all lines	Top Hit Database Source		HOMAN	Т	Т	NOM		NAMI IL FOR	HUMAN	Τ				EST_HUMAN	HUMAN	LN	NT	TN	EST HUMAN	TIVE TIVE	2 2	N _T	LN		T L	LV	EN .	EST HUMAN		EST HUMAN	
All C	Top Hit Acesslon No.		-		T			T	AW 898080.1	24.7	143021		D84122.1	AW583858.1	AW583858.1	11427788 NT	1.0E-121 AF064200.1	30334		1.0E-121 AU119320.1	TN 11520170170	1.0E-122 At 1.150.176 NT	A E 1 1 4 4	1.0E-122 N. 1.1.200.	1.0E-122 M20707.1	22/AF 107 700.1		900	22 BEBUOUZ4.	22 BF3161 /0.1	- Colored
	<u> </u>	\ aine	1.0E-121 A		_								1.0E-121	1.0E-121		1.0E-121	1.0E-121	1.0E-121	1.0E-121 N59624.1	1.0E-121	1.0E-122	١			1.0E-1	1.0E-1	1	1.01-1	1.0E-1	1.0E-1	1.05-1
	Expression Signal	-	2	3.24	0.97	0.85	16.0	0.68	1.78	1.78	2.11	2.45	2.45	1.21	1.21	2.95	1.28	3.46	2.53	2.83	2.28	2.63	2.14	3.88							5.21
	ORF SEQ ID NO:		29863	30259	30510	31167	32294			30538		33664	33665	35615		1	36848		١	3 37269				1 26302	0 28614	2 27128		0 27151		11	2 27955
	Exan SEQ ID	<u>_</u>	17035	17847	17994		L	1_	17943	17943	20533	20537	1_	<u> </u>	I	Į.	l	23569	上	L	L	7 13128		2 13631	13950	ļ	7 14450	14450	14547	1 1	15212
	- 0		A20A	4010	5186	5474	6524	6788	9889	8888	7838	7842	7842	9758	07.60	10875		10683	10917	11309	284	327	348	862	1198	1688	1707	1707	1807	2495	2495

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Siligia Extil Flobes Explassed III Didili	Top Hit Descriptor	Homo saplens FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2 mRNA, complete cds	Homo sapiens amyloid beta (A4) precursor protein (protease nextn-ll, Abheimer disease) (APP), mRNA	UI-HF-BN0-all-a-03-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3079948 61	601113567F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3354232 6	601113567F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3354232 6	ak49h06.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1409339 3	Homo sapiens mRNA for doublesex and mab-3 related transcription factor 1 (UMKL1)	Homo sapiens lethal giant larvae (Drosophila) homolog 2 (LLGL2), mKNA	lg/s2h07.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2013757 3' similar to sw:wTA1_HUMAN Q13330 METASTASIS-ASSOCIATED PROTEIN MTA1.;	qy32h07.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:2013767 3' similar to sw.mTa1_HUMAN Q13330 METASTASIS-ASSOCIATED PROTEIN MTA1.:	Novel human gene mapping to chomosome X, isoform of dbi (proto-oncogene)	EST367904 MAGE resequences, MAGD Homo sapiens cDNA	Homo saplens gene for B120, exon 10	Homo saplens phosphomannomulase 1 (PMM1), mRNA	802018058F1 NCI_CGAP_Brn67 Homo saplens cDNA clone IMAGE:4153670 5	802018058F1 NCI_CGAP_Brn67 Homo sapiens cDNA clone IMAGE:4163670 5	Hamo saplens chromosome 21 segment HS21C049	Homo sapiens inner membrane protein, mitochondrial (mitofilin) (iMMT), mKNA	Homo sapiens phosphatidylinositol-4-phosphate 6-kinase, type II, beta (PiP6K2B) mRNA, and translated products	Homo sapiens phosphatidylinositol-4-phosphate 5-kinase, type II, beta (PIP5K2B) mRNA, and translated products	Homo saptens partial mRNA for immunoglobulin kappa chain variable region (IGVK gene), sample GN02	Human amelogenin (AMELY) gene, 3' end of cds	Human amelogenin (AMELY) gene, 3' end of cds	Human amelogenin (AMELY) gene, 3' end of cds	Homo saplens RAB9-like protein (LOC51209), mRNA	Homo saplens glutaminyl-peptide cyclotransferase (glutaminyl cyclase) (QPCT), mRNA	Homo sapiens retinaldehyde-binding protein (CRALBP) gene, complete cas
DI L IOY DIE	Top Hit Database Source	ħ	ŇŢ	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	N-	M	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	N	LΝ	EST_HUMAN	EST_HUMAN	NT	NT	ΤΝ	Z	LV.	۲	NT	FN	ΝΤ	TN	ΝΤ
0	Top Hit Acession No.	1.0E-122 AF264717.1	4502168 NJ	\W504645.1	1.0E-122 BE256039.1	3E256039.1	1.0E-122 AA868671.1	1.0E-122 AJ276801.1	11424216 NT	1.0E-122 Al359618.1			1.0E-122 AW955834.1	22 AB024068.1	11418187 NT	3F345274.1	1.0E-123 BF345274.1	1.0E-123 AL163249.2	5803114 NT	4505818 NT	4505818 NT	1 0E-123 A.1388641.1	M55419.1	M55419.1	M55419.1	7705962 NT	6912617 NT	L34219.1
	Most Similar (Top) Hit BLAST E Value	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1 0F-123	1.0E-123 M55419.1	1.0E-123 M55419.1	1.0E-123 M55419.1	1.0E-123	1.0E-123 6	1.0E-123
	Expression Signal	1.41	5.04	1.46	1.36	7.1	6.73	0.65	1.21	1.19	1.19	1.05	2.17	1.88	9.9	1.74	1.74	5.4	2.5	5.58		 	2.7	2.7	2.7	3.62	0.95	1.56
	ORF SEQ ID NO:	28260	30148	L	31170	31170	32865		34761	1						L	26182		L	<u> </u>				27558			28657	
	Exan SEQ ID NO:	15612	17526	17858	18275	18275	19801	21387	21617	21902	21902	22682	23588	24046	24509	13523	13523	13754	13761	13968	13966	14185	14823	14823	14823	15038	16007	18163
	Probe SEQ ID NO:	2844	4795	4930	5478	8999	7113	8695	8928	9223	9223	10034	10908	11358	11958	751	761	992	1001	1218	12.4	2 2	360	2002	2000	23.43	3245	5361

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Top Hit Descriptor	Homo sapiens retinaldehyde-binding protein (CRALBP) gene, complete cds	601591108F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3845433 5'	AU118435 HEMBA1 Homo sapiens cDNA clone HEMBA1003591 5'	yq84a03.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:202444 5' similar to SP:YAK1_YEAST P14680 PROTEIN KINASE YAK1;	Human growth hormone releasing hormone gene, exon 7	Human hBRAVO/Nr-CAM precursor (hBRAVO/Nr-CAM) gene, complete cds	Homo sapiens heparan sulfate (glucosamine) 3-O-sulfotransferase 2 (HS3ST2), mRNA	Homo sapiens 2'-6'digoadenylate synthetase 2 (OAS2), mRNA	601162815F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3509162 5'	AU131881 NT2RP3 Homo sapiens cDNA clone NT2RP3003409 5'	AU131881 NTZRP3 Hamo sapiens cDNA clone NT2RP3003409 5'	RC4-BT0311-251199-012-e07 BT0311 Homo saplens cDNA	Homo sapiens mRNA for KIAA0454 protein, partial cds	Oryctolagus cuniculus New Zealand white elongation factor 1 alpha (Rabefla2) mRNA, complete cds	602086791F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4250879 5'	602086791F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4250879 57	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens DNA for amylold precursor protein, complete cds	Homo sapiens chromosome 21 segment HS21C046	z/81b04.rl Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:728719 6' similar to TR:G300482 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) ;	zt81b04.r1 Stratagene schizo brain S11 Homo sapiens cDNA done IMAGE:728719 6' similar to TR:G300482 G300482 POL-REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) ;	Human putative ribosomal protein S1 mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens hypothetical protein (HSPC068), mRNA	Homo sapiens glucose transporter 3 gene, exons 9, 10, and complete cds	Homo sapiens glucose transporter 3 gene, exons 9, 10, and complete cds	Homo sapiens mRNA for nucleolar RNA-helicase (noH61 gene)	601491715F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3893954 6	Homo saplens glutamate receptor, ionotropic, kainate 1 (GRIK1) mRNA
Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	FN	LN	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN LN	NT	EST_HUMAN	EST_HUMAN	IN	LN.	TN	LN	EST_HUMAN	EST_HUMAN	- LN	K	ĮN.	FX	ΤN	TN	EST_HUMAN	٦
Top Hit Acession No.	1.0E-123 L34219.1	3E799746.1	1.0E-123 AU118435.1	153198.1	J42224.1	J55258.1	11525833 NT	11436439 NT		1.0E-123 AU131881.1	1.0E-123 AU131881.1	4W371924.1	1.0E-123 AB007923.1	109823.1	1.0E-123 BF677292.1	1.0E-123 BF677292.1	4507500 NT	4507500 NT	087675.1	24 AL163246.2	1.0E-124 AA397551.1	1.0E-124 AA397551.1	1.0E-124 AF155654.1	4507500 NT	7705446 NT		1.0E-124 AF274892.1	1.0E-124 AJ131712.1	1.0E-124 BE879524.1	4504116 NT
Most Similar (Top) Hit BLAST E Value	1.0E-123	1.0E-123	1.0E-123	1.0E-123 H53198.1	1.0E-123 U42224.1	1.0E-123 U55258.1	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123	1.0E-123 U09823.1	1.0E-123	1.0E-123	1.0E-124	1.0E-124	1.0E-124 D87675.1	1.0E-124	1.0E-124	1.0E-124	1.0E-124 /	1.0E-124	1.0E-124	1.0E-124	1.0E-124 /	1.0E-124	1.0E-124	1.0E-124
Expression Signal	1.56	1.62	2.59	1.2	1.25	2.87	1.62	1.3	2.18	79'0	79.0	1.13	2.43	15.48	4.66	4.66	2.19	2.19	2.99	2.84	2.68	2.68	7.84	1.61	1.94	4.95	4.95	2.29	3.05	0.85
ORF SEQ ID NO:			32145	32688	32698	32849	33061	33312	33324	33636	33637		35193	35239	37637	37638	26708	25709		25898	26092	26093	26174	26223	26321	26747	26748	27263	27512	28774
Exan SEQ ID NO:	i i		19146	19643	19652	19783	19985	20212		<u>L.</u>	20511		22023	22067		24314	13070	13070	13076	13259	13450	13460	13515	13562	13653	L	14074	ı	1	16118
Probe SEQ ID NO:	5361	5494	6377	6905	6915	7094	7302	7542	7551	7816	7816	8433	9269	9405	11720	11720	292	262	268	473	675	675	742	790	88	1326	1325	1808	2054	3358

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Homo sapiens ubiquitin specific protease 9, X chromosome (Drosophila fat facets related) (USP9X), mRNA hg94a09.x1 NCI_CGAP_Kid11 Homo sepiens cDNA done IMAGE:2953240 3' similar to TR:095162 095162 PEROXISOMAL SHORT-CHAIN ALCOHOL DEHYDROGENASE.; hg94e09.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2953240 3' similar to TR:095162 095162 PEROXISOMAL SHORT-CHAIN ALCOHOL DEHYDROGENASE; Homo sapiens ATP-sensitive inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, exon ac08h05.s1 Stratagene HeLa cell s3 937216 Homo saplens cDNA clone IMAGE:866897 3 UI-HF-BN0-akz-b-04-0-UI.r1 NIH_MGC_60 Homo sapiens cDNA clone IMAGE:3078846 6' Homo sapiens ATP-sensitive Inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, H.sapiens lactate dehydrogenase B gene exon 1 and 2 (EC 1.1.1.27) (and Joined CDS) Human muscle glycogen phosphorylase (PYGM) gene, exons 6 through 17 hj05c06.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2980906 3 Homo sepiens IQ motif containing GTPase activating protein 1 (IQGAP1) mRNA 602124644F1 NIH_MGC_66 Homo sepiens cDNA clone IMAGE:4281636 5¹ AV711283 Cu Homo sepiens cDNA clone CuAADF07 5 wc43g03.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2321428 3' wc43g03.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone IMAGE:2321428 3' AV645633 GLC Homo sapiens cDNA done GLCACE04 3' CGAP_Kid12 Homo sepiens cDNA clone IMAGE:2400891 3 Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA 600943771F1 NIH_MGC_8 Homo saplens cDNA clone IMAGE 2966585 5 600943771F1 NIH_MGC_8 Homo saplens cDNA clone IMAGE:2866585 5 Homo sapiens cep250 centrosome associated protein mRNA, complete cds Homo saplens cep250 centrosome associated protein mRNA, complete cds w83f02.x1 NCI_CGAP_Kid12 Homo saplens cDNA clone IMAGE:2400891 Homo seplens gluternate receptor, ionotropic, kainate 1 (GRIK1) mRNA Homo sepiens gene for B120, exon 11 Human fibronectin gene extra type III repeat (EDII), exon x+1 Homo sepiens hypothetical protein FLJ10300 (FLJ10300), mRNA Top Hit Descriptor Homo sapiens leucine-rich, glioma inactivated 1 (LGI1), mRNA AV645633 GLC Homo sepiens cDNA clone GLCACE04 3 Homo saplens ribosomal protein L5 (RPL5) mRNA M.musculus mRNA for hoxe3 gene Single Exon Probes Expressed in Brain wi93f02.x1 NCI Top Hit Database HOMAN EST HUMAN HUMAN HUMAN HOMAN EST HUMAN HUMAN **EST_HUMAN** HUMAN **EST HUMAN** Source EST_HUMAN EST HUMAN HUMAN EST HUMAN HUMAN 몂 NT EST 4504116 NT 11420654 NT 4506654 N 눋 Top Hit Acession 4507500 8922337 4506786 BE271296.1 BE271295.1 1.0E-124 AW612106.1 BF696135.1 1.0E-124 AB024069.1 1.0E-124 AV711263.1 AA630331.1 AW612106.1 Š .0E-124 AI799864.1 1.0E-124 S78684.1 1.0E-124 S78684.1 1.0E-124 AI799864.1 1.0E-124 AV645633. .0E-124 AV645633. 1.0E-124 AF022655. .0E-124 AI767133.1 1.0E-124 AI767133.1 1.0E-124 AF022655. 1.0E-124 X13794. **U94776.1** 1.0E-124 AW 503 1.0E-124 1.0E-124 1.0E-124 1.0E-124 E 1.0E-124 (Top) Hit BLAST E Most Simila 1.0E-124 1,0E-124 1.0E-124 1.0E-124 12.12 0.92 <u>2</u> 6.94 0.98 0.94 1.26 0.61 0.62 2.31 0.62 7.57 0.91 1.26 Expression 8.07 1.57 3.51 1.61 Signal 29033 29269 29432 30061 31514 ORF SEQ 31293 30647 32099 32695 32786 32787 34195 33982 34196 35310 ÖΝΟ 35342 36905 35630 37265 SEQ ID 16239 16393 18383 **16630** 18023 18588 19109 19649 22062 18856 19729 19729 21065 22062 20128 20850 22130 1680 22161 23487 23964 Probe SEQ ID 3482 3482 3840 4056 4896 4881 5215 5586 8912 6339 8362 5797 6077 7337 7452 8362 0908 9390 8390 9508 8156 9771 10804 11305 9477 9477 9508 10976

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z81b04.r1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:728719 6' similiar to TR:G300482 G300482 POL≃REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) ; nv59a08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3177686 3' similar to TR:Q25058 Q25058 nv59a08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3177686 3' similar to TR:Q25058 Q25058 \$19e03.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2141980 3' similar to TR:031662 031662 119e03 x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2141980 3' similar to TR:031662 031662 zt81b04.r1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:728719 5' similar to TR:G30048 Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein 2633c07.s1 Scares, pregnant, uterus, NbHPU Homo sapiens cDNA clone IMAGE:488540 3' similar to gb:X65857, cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN); 263-07.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:486540 3' similar to gb:X65857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN); zi01g09.r1 Soares fetal liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:429588 5 G300482 POL=REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT); Homo sepiens KIAA0744 gene product, histone deacetylase 7 (KIAA0744), mRNA Homo sepiens KIAA0022 gene product (KIAA0022), mRNA Homo sapiens mRNA for KiAA1172 protein, partial cds 601577981F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3926686 5 Homo sepiens calcineurin binding protein 1 (KIAA0330), mRNA Homo sepiens calcineurin binding protein 1 (KIAA0330), mRNA Top Hit Descriptor HA0086 Human fetal liver cDNA library Homo sapiens cDNA HA0086 Human fetal liver cDNA library Homo sapiens cDNA Homo sapiens mRNA for KIAA1093 protein, partial cds Homo saplens Usurpin-alpha mRNA, complete cds Homo sapiens chromosome 21 segment HS21C010 fomo saplens Usurpin-alpha mRNA, complete cds Homo sapiens ALR-like protein mRNA, partial cds (L44L) and FTP3 (FTP3) genes, complete cds Homo sapiens inhibin, alpha (INHA) mRNA Homo sapiens Inhibin, alpha (INHA) mRNA YKRS PROTEIN.; FIBROPELLIN IA YKRS PROTEIN. FIBROPELLIN IA **EST HUMAN** EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN HUMAN EST_HUMAN **EST_HUMAN** EST_HUMAN Top Hit Database Source EST 11417862 NT 11417862 NT 4504696 NT 4504696 NT 눈 눋 눋 눋 7662279 N **Top Hit Acession** 1.0E-125 AA042813.1 1.0E-125 AL163210.2 1.0E-125 BE219510.1 1.0E-125 AA042813.1 1.0E-124 AA397551.1 1.0E-124 AA397551.1 1.0E-124 AB029016.1 1.0E-125 BE219510.1 1.0E-125 AB032998.1 1.0E-125 AI110656.1 AF264750.1 1.0E-125 AF015450.1 .0E-125 AF015450.1 1.0E-125 AA011278.1 1.0E-124 AI446455.1 1.0E-124 AI446455.1 1.0E-125 AI110656.1 U78027.1 1.0E-125/ .0E-125 1.0E-125 1.0E-125 1.0E-125 1.0E-125 1.0E-124 1.0E-125 (Top) Hit BLAST E Most Similar Value 1.74 1.18 1.76 4 4. <u>‡</u> 1.74 5.49 87 1.59 0.99 1.3 6 5.47 0.99 9.9 1.68 Expression Signal ORF SEQ ID NO: 26405 36452 36453 26092 26093 31026 30729 25603 25604 26043 26044 26134 26545 27242 27259 27816 27967 28057 28058 25441 30730 13408 13485 13610 23219 23219 13450 13450 13115 12828 13408 15225 SEQ ID 24823 12961 12961 13887 14533 14544 15317 14544 15317 15080 1557 ģ 978 2604 2604 2604 Probe SEQ ID 11452 11452 12029 12029 12706 629 711 1131 1793 2358 12454 <u>8</u> 46 311 629 1804 1804 417 1668 ġ

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	Top Hit Descriptor	bb74f06.y1 NIH_MGC_12 Homo saplens cDNA clone IMAGE:3048131 5' similar to TR:095604 095504 ZINC FINGER PROTEIN:	zk83c07.s1 Soares_pregnant_uterus_NDHPU Homo saptens curin curin linnstillatore curin gb.X68857_cds1 OLFACTORY RECEPTION-LIKE PROTEIN HGMPO7E (HUMAN);	Homo sapiens zinc finger protein ZNF287 (ZNF267), minus	Homo saplens zinc finger protein ZNr28/ (ZNr28/), mrNvx Homo saplens zinc finger protein ZNr28/ (ZNr28/), mrNvx	M69808.X1 NCT CCAP_LUZ4 natio september con Notice in the Company of the PIBROPELLIN IA: PIBROPELLIN IA: PERCAPA A ICT CCAP 1.124 Home september CDNA clone IMAGE:3177886 3' similar to TR:Q25058 Q25058	FIBROPELLIN IS.	Homo sapiens Kildausoo piotein (Nichaesoo), mistro	QV2-HT05/7-010900-163-906 F103/7 HQIIO 84/1616 OCHA	6014334/2F1 NIT MICC /2 FULL Sapiens CONA clone IMAGE:2256108 3' similar to WP:C45G9.2	CEOURSE : CARL CARL CARL CARL CARL CARL CARLO CA	601305870F1 NIH MGC 39 Home saplens CDNA clone IMAGE:3689790 5	0013309207 I Nin JWO 44 Home saniers CDNA clone IMAGE:3889790:5'	0015556207 I NIT MCC_44 I NIT CAPACITO CONTROL	TONIO Bapicale TO - il gono, con i	ROLLO BAPTELS TOTAL BATTS, WALLE SADIENS CON CONE IMAGE:3608084 5		Human chromosome 10 duplicated adrenoleukodystrophy (ALD) gene segment containing exons 8-10	Human chromosome 10 duplicated adrenoleukodystrophy (ALD) gene segment containing exons 8-10	QV1-H10838-0/0600-191-d12 H10839 Humo satisfies CDNA	CAVI-H 10038-0/1000-191-01-1 Homo saniens cDNA clone IMAGE:2171981 3' similar to TR:Q14089 Q14089	INSZEDS.XI INCLOCAL JAMAT I SOMO CAPATOR I SOMO SEDENTIAL MYPOTETTICAL PROTEIN SEDIENS CDNA clone IMAGE:3944631 5'	601 690 340F1 Nir MGC / Troins septents controlled to the man ment of free transfer of the MARSON came, partial cds	Human Introduce and B	Homo saniens ryanodine receptor 1 (skeletal) (RYR1), mRNA	Homo saplens mRNA for KIAA0697 protein, partial cds	
ביוומוס ביעמון ו	Top Hit Database Source	EST_HUMAN 2	M3.1 EST_HUMAN			EST_HUMAN	T_HUMAN		П	EST_HUMAN		\neg	٦	HOMAN	Į.	Т	ESI HOMAIN	Ę	TN	EST_HUMAN	EST HOMAN		EST HUMAN	N.	Z	LN LN	, NI
27	Top Hit Acession No.	1.0E-125 BE018009.1	25 AA042813.1	11425114	11425114 NT	25 BE219510.1	25 BE219510.1	11436448 NT	25 BE175169.1	1.0E-125 BE892660.1	25 AI679904.1	1.0E-125 BE736055.1	26 BE562526.1	125 BE562526.1	25 X03427.1	125 X03427.1	25 BE515100.1	25 U90288.1	125 U90288.1	125 BE181640.1	25 BE181640.1	125 A1565996.1	125 BE794576.1	125 AB002298.1	AF0434	125 14255/0 NI	AB014507.1
	Most Similar (Top) Hit BLAST E Value	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1.0E-126	1.0E-125	1.0E-126	1.0E-125	1.0E-1	1.0E-125	1.0E-125	1.0E-1	1.0Ē	1.0E-1	1.0E-1	1.0E-	1.0E-	1.0E	1.0	8	9
	Expression Signal	0.94	0.92	2.09	2,09	1.48	1.48	3.16	0.91	3.76	0.74	8.0	1.63	1.53	5.26	5.26	0.55	0.99		6.83							3.94
	ORF SEQ ID NO:	28425		ļ	2984	<u> </u>	25604				31611						33483	34273	34274			35207	L	36253	Ц		1 37028
	Exon SEQ ID NO:	17874	1	ı	l	<u> </u>	12961	1_		١_	1	1_	L		L	L	20370	21136	1		1	ſ	1_	<u> </u>			1 23751
	Probe SEQ ID NO:	0008	SONS C	3000	4513	4971	4971	5783	5802	5842	5884	6188	8488	9486	6961	6961	7706	8444	0,444	9018	9018	9281	1885	1039	10581	10758	11081

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	Top Hit Descriptor	Lame caniens myosin, heavy polypeptide 1, skeletal muscle, adult (MYH1), mRNA	Homo saniens poly(A) binding protein II (PABP2) gene, complete cds	IRC3-ST0186-250200-018-c11 ST0186 Homo saplens cDNA	10V3-BT0569-020200-075-009 BT0569 Homo saplens cDNA	QV3-BT0569-020200-075-g09 BT0569 Homo saplens cDNA	Homo sapiens CDC-like kinase (CLK) mRNA	H saplens gene for alphat-antichymotrypsin, exon 3	Homo sapiens hypothetical protein FLJ20048 (FLJ20048), mRNA	Homo sapiens hypothetical protein FL J20048 (FL J20048), mRNA	Homo saplens RAN binding protein 2 (RANBP2), mRNA	Homo saniens glutamate receptor, ionotropic, kalnate 1 (GRIK1) mRNA	3722013 dt Stratagene pangreas (#637208) Homo saplens cDNA clone IMAGE:392420 3	2722013 11 Stratagene pancreas (#937208) Homo sapiens cDNA clone IMAGE:592420 5	H saniens DNA for liver cytochrome b5 pseudogene	Homo septent death receptor 6 (DR6), mRNA	_	7	_	1	$\neg T$	Homo saprens mRNA for KIAA1525 protein, partial cds	Home seriens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cds	Homo saplens ciliary dynein heavy chain 9 (DNAH9) mRNA, complete cdis	Т	Т	\neg	Homo saplens minny for Nink 1224 process, process	Homo sapiens might for the form of the for	Human mRNA for ankyrin (variant 2.1)	HUMAN GEROOBS Similar to SW:TSG6_HUMAN			
Si i liovi eigilio	Top Hit Database Source		Z	New In	TOT CLIMANI	EST HIMAN	NT INDING	Z I	E	F14	LIV.	į į	NAME OF THE PARTY	TOT LOUIS	NICHOLINE THE	Z	LANNI LI	TOT TOWN	TOTAL TOTAL	EST HOMEN	EST HUMAN	5	Z	Z	NAME OF THE PARK		EST_HUMAN	TN	L'N	FN		FST HUMAN	NT NT	
	Top Hit Acession No.		39505			T	E074257.1	4/5800/ IN	58735.1 COMOGE NT	882000	FIN 929290	1 N 0 70 70 70 1	4504115 NI	1.0E-126 AA160709.1	1.0E-126 AA160709.1	(53941.1	20/09/	1	2.1	T66998.1	26 AA460075.1	26 AB040958.1	26 AB040958.1	26 AF257737.1	126 AF257737.1	26 AU136463.1	26 A1806483.1	126 AB037715.1	26 AB037715.1	26 X16609.1		A A 402368 1	4505424 NT	
.	<u>a</u> + 111	Value	1.0E-125	1.0E-125	1.0E-125 A	1.0E-125 BE074267.1	1.0E-125 BE074267.1	1.0E-126	1.0E-126 X68735.1	1.0E-126	1.0E-126	1.0E-126	1.0E-126	1.0E-128/	1.0E-126/	1.0E-126 X53941.1	1.0E-126	1.0E-126 N34078.1	1.0E-126	1.0E-126 T66998.1	1.0E-126	1.0E-126	1.0E-126	1.0E-126	1.0E-126	1.0E-126	1.0E-	106	1 0F-1	1 0E-1		,		
	Expression Signal .		1.74	4.84	1.92	3.58	3.58	1.46	1.45	1.17	1.17	1.48	0.72	7.54	7.54	1.09	1.6	1.74	0.81	0.68	3.22	4.2	4.2			0.62	69 0						0.99	
	ORF SEQ E		37213	37218	37284	37397	37398	26189	26331	27804	27805	28059	-	28479			29040	İ	30413	31325	31887				33158	33375	90,400	1			2			35539
			23921	23927	23984	24087	24087	13529	13667	15067	L	15318	L	L	L	<u> </u>	L	١.	L	1	18017		L	1_	1_	L	_	1	-1		7 20582	 -	1	22346
		ë	11259	11265	11377	11486	11486	757	899	2344	2344	2605	3069	3070	3070	3620	3847	4783	5078	5616	00.70	8107	6107	7399	7399	7602		7655	7778	7778	7887		8083	9698

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01/57275	-	Γ		<u>-</u>	_	Γ-	ΓT					_	-	_			_	 -	_	 -	-	<u></u>	-	P(CT.	/U	501	/00
Top Hit Descriptor	[-] Hardin management	Primital Hacrophiage marriose receptor (MRC1) gene, expn 5	NOZ.139136F1 NIH_MISC_46 Home sapiens cDNA clone IMAGE:4298240 6	R0157708154 NILL MCC 611	HOUSE STATE THAT MICE BY HOME SEPTENDED TO A CIONE IMAGE:3926685 5'	Home serious mb/A for casen Knase I epsilon, complete cds	Homo sanlens mRNA for caselir kinnes I epsilon, complete cds	Homo sapiens mBNA for casalit Vienna Landing.	Homo sablens DNA for amyloid predictor protols Amelois Amelois	Homo saplens DNA for amyloid pregired protein complete de	Homo saplens intersectin short isoform (TTSN) mBNA Annuals and	Homo saplens lost on transformation I OT1 mBNA commissions	Homo saplens ublautifu specific professe 8 /1 ISD81 with A	Homo saplens leukocyte Immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), mRNA	Homo sapiens leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), mBNA	Children and and and and and and and and and an	Home sections all the section of the	Himes month for many complete cds	rx428/29 1 Sparse fore form NEOLES A	2x42a02.11 Soares total faths Nh2HE8 ou Home capiens cDNA clone IMAGE:788098 6'	au80e06.y1 Schneider felal brain 00004 Homo sapiens cDNA clone IMAGE:2782594 6' similar to TRANSCRIPTION FACTOR S-II-RELATED PROTEIN ;contains element MER22	repetitive element	rionno sapiens delayed rectifier potassium channel subunit IsK mRNA, complete cds	romo sapiens neuroblascome-amplified protein (LOC51694), mRNA	Horno sapiens neurobiastoma-amplified protein (LOC51694), mRNA	Home capiens BAD4 / Complete cds	Homo sapiens chromosome 24	Homo sapiens Ringt and YY4 hindran ander, PSPBD
Top Hit Database Source	FZ	EST LIBRARI	FST HIMAN	EST HIMAN	TW.	LZ	Į.	LN	NT	NT	LN	NT.	FX	Ę					T HIMAN	HUMAN		EST HOMAN						
Top Hit Acession No.	M93198 1	BF883175 1	1.0E-126 BE261660.1	1.0E-126 BE743922 1	Γ						_		4827053 NT	- 5803065 NT	5803085 NT	4506620 NT	246505.1						06030	TIM OCCOUNT	F252207 4 IN	6384	16326	6912639
Most Similar (Top) Hit BLAST E Value	1.0E-128 M	1.0E-128 R	1.0E-126	1.0E-128	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127 D87675.1	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127 AF	1.0E-127 X12881 1	1.0E-127 AA450131.1	1.0E-127 AA450131.1	4 0E 407 AM464007 4	1.0E-127 AF135188 4	1 OF 127	1 0E-127	1.0F-127 AF252207 4	1.0E-127	1.0E-127 AL	1.0E-127
Expression Signal	1.27	2.38	6.47	7.17	8.4	4.63	4.71	4.71	3.54	3.54	2.03	1.37	1.08	2.44	2.44	6.02	2.73	3.04	1.1	1.1	7	0.7	23.74	23.74	0.94	4.35	1,92	1.26
ORF SEQ ID NO:	36624	36692	37414	30413	26822	25623	25622	25623	25716	25717	26301	28330	27128	27515	27516	27661	27801	28068	28081	28082	29178	29465	29592	29593	29824	28926		29933
SEQ ID	23384	23450	24102	17797	12982	12982	12982	12982	13075	13076	388	13664	54 88	14790	14790	14926	15064	15325	15338	15338	16543	16838	16969	16968	17198	17299	17325	17360
Probe SEQ ID NO:	10693	10766	11501	12490	168	168	169	169	287	267	861	88	1888	2058	2058	2197	2341	2614	2826	2626	3791	4096	4227	4227	4462	4564		4825

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		_	_						_						_ *	*	~	н		Jad*	400	_	4				-	arar ranchi	·bad
Top Hit Descriptor	za01a10.r1 Soares melancoyte 2NbHM Homo sapiens cDNA clone IMAGE:291268 5' similar to SW:PiPe_RAT P10688 1-PHOSPHATIDYLINOSITOL-4, 5-BISPHOSPHATE PHOSPHODIESTERASE DELTA 1;	Homo sapiens neuronal cell adhesion molecule (NRCAM) mRNA	H.sapiens NOS2 gene, exon 6	H.sapiens TCF11 gene, exon 3-6	Homo sapiens Integrin, beta 8 (ITGB8) mRNA	Homo saplens Immunoglobulin superfamily, member 3 (IGSF3), mRNA	Homo sapiens reelin (RELN) mRNA	Homo saplens Pendred syndrome (PDS), mRNA	Homo sapiens Pendred syndrome (PDS), mRNA	QV3-BN0046-150300-121-h11 BN0046 Homo saplens cDNA	Homo sapiens Chediak-Higashi syndrome 1 (CHS1), mRNA	Homo sapiens Chediak-Higashi syndrome 1 (CHS1), mRNA	Homo sapiens secretory pathway component Sec31B-1 mRNA, alternatively spliced, complete cds	Homo sapiens secretary pathway component Sec31B-1 mRNA, alternatively spliced, complete cds	qm94h09.x1 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1896449 3'	Homo sapiens Chediak-Higashi syndrome 1 (CHS1), mRNA	Homo sapiens similar to heat shock 70kD protein 9B (mortalin-2) (H. sapiens) (LOC63184), mRNA	Homo sapiens similar to heat shock 70kD protein 9B (mortalin-2) (H. sapiens) (LOC63184), mRNA	601434784F1 NIH_MGC_72 Hamo sapiens cDNA clane IMAGE:3919917 5'	601434784F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3919917 5	Homo sapiens mRNA for casein kinase I epsilon, complete cds	Homo sapiens mRNA for casein kinase I epsilon, complete cds	Homo sapiens gene for AF-6, complete cds	601278127F1 NIH_MGC_20 Hamo sapiens cDNA clone IMAGE:3618822 5'	Human FAU1P pseudogene, trinucleotide repeat regions	Human FAU1P pseudogene, trinucleotide repeat regions	Homo saplens ribosomal protein S2 (RPS2) mRNA	Homo sapiens chromatin-specific transcription eiongation factor, 140 kDa subunit (FACTP140), mRNA	Homo saplens mRNA for KIAA1247 protein, partial cds
Top Hit Database Source	EST_HUMAN	LN	LN	TN	IN	TN	LN	ΙN	TN	EST_HUMAN	N	TN	TN	LΝ	EST HUMAN	TN	NT	NT	EST_HUMAN	EST HUMAN	NT	LN	NT	EST_HUMAN	TN	LN	NT		NT
Top Hit Acession No.	W03547.1	4826863 NT	X85764.1	X84060.1	4504778 NT	11421595 NT	4826977 NT	11421914 NT	11421914 NT	AW996292.1	11427235 NT	11427235 NT	AF274863.1	AF274863.1	AI298932.1	11427235 NT	11417339 NT	11417339 NT	BE895415.1	BE895415.1	AB024597.1	AB024597.1	AB011399.1	BE385617.1	U02523.1	U02523.1	4508718 NT	11437455 NT	AB033073.1
Most Similar (Top) Hit BLAST E Vælue	_	1.0E-127		1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127		1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	_		1.0E-127	,	1.0E-127		1.0E-128		_	1.0E-128	1.0E-128	1.0E-128
Expression Signal	1.37	2.4	4.25	2.17	5.28	0.89	0.81	1.65	1.65	0.64	9.0	0.8	4.17	4.17	-	1.34	7.88	7.88	3.25	3.25	2.25	2.25	2.88	3.04	5.5	5.5	8.76	1.1	1.08
ORF SEQ ID NO:	31329	31359	31428	31813	31979		32461	33461	33462	33470	34625	34626	35377	35378	35631	36101	37050	37051	37548	37547	25622	25623	31048	25873	. 27520	27521	27672		28802
Exan SEQ ID NO:	18416	18445	18506	18849	19003		19444	20348	20348	20355	21477	21477	22193	22193	22424	22889	23777	23777	24224	24224	12982	12982	24814	13233	14795	14795	14934	15165	16148
Probe SEQ ID NO:	5620	5650	5713	6070	6229	0999	6962	7684	7684	7697	8785	8785	9540	9540	8773	10241	11107	11107	11627	11627	12244	12244	12444	447	2063	2063	2206	2446	3389

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Top Hit Descriptor	Homo sapiens prospero-related homeobox 1 (PROX1), mRNA	H. sapiens gene for Inter-alpha-trypsin inhibitor heavy chain H1, exon 12	601580466F1 NIH MGC 9 Homo saplens cUNA clone IMAGE: 3929057 5	601580466F1 NIH_MGC_9 Homo sapiens curva cione invade: 3929037 o	Homo sapiens phosphodiesterase 1C, calmodulin-dependent (10kU) (10C1C), month	7q86b10.x1 NCI_CGAP_Lu24 Homo sapiens cunA cigne invage: 3	601503846F1 NIH MGC 71 Homo sapiens CUNA cione invade: 3803784 3	Homo sapiens putative ABC transporter (VVIII EZ), micros	Homo saplens mRNA tor KIAA0454 protein, parda cos	Homo sapiens mRNA for KIAAU454 protein, parael cas	ns04a11.1 NCI_CGAP_EWI Home septens cUNA done invace: 1 102020 similar to Inc. 5201.00 COLTAGE CHROMOSOME SEGREGATION GENE HOMOLOG CAS.;	Homo sapiens glutamate receptor, Ionotropic, N-methyl D-asparate 2D (GRINZD), mrinn	om68h08.s1 NCI_CGAP_GC4 Homo saplens cDNA clone IMAGE:1652383 3' similar to ge:X64841 C7 CLINA DEPENDENT KINASES REGULATORY SUBUNIT 1 (HUMAN);	Homo sapiens mRNA for TRABID protein (TRABID gene)	601277826F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE:3818750 5'	EST367360 MAGE resequences, MAGC Hamo sapiens cDNA	insulin-like growth factor binding protein-2 [human, placenta, Genomic, 1019 nt, segment 2 of 4]	insulin-like growth factor binding protein-2 [human, placenta, Genomic, 1019 nt, segment 2 of 4]	Novel human mRNA containing Zinc finger C2H2 type domains	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	genes, conjugac dus superiores S. transferase theta 2 (GSTT2) and clutathione S-transferase theta 1 (GSTT1)	genes, complete cds	Homo saplens zinc finger protein 76 (expressed in testis) (ZNF78), mRNA	ZINC FINGER PROTEIN HZF10	ZINC FINGER PROTEIN HZF10	ZINC FINGER PROTEIN HZF10	Homo sapiens mRNA for KIAA1459 protein, partial cds	CM/YA5 Human cardiac muscle expression library Homo sapiens cDNA clone 4151935 similar to CMYA5	Cardiomyopathy associated gene 5	CMYA6 Human cardiac muscle expression library Homo sapiens cUNA cione 4151835 similar to CM1A5 Cardiomyopathy essociated gene 5	
Top Hit Database Source			HUMAN	T HUMAN	Z.	EST_HUMAN	EST HUMAN	N ₁	LN-	NT L	EST_HUMAN	۲	EST HUMAN	N	EST_HUMAN	EST HUMAN	TN	LN	۲		Ž	Į.	۲	SWISSPROT	SWISSPROT	SWISSPROT	_ - 		EST_HUMAN	EST_HUMAN	
Top Hit Acessian No.	11426673 NT	X69539.1	BE747981.1	BE747981.1	11420965 NT			15923	AB007923.1	AB007923.1	AA639198.1 EST	11425264	A A 926959.1	1.0E-128 A.1262060.1		L		337722.1	1.0E-129 AL096880.1		1.0E-129 AF240786.1	1.0E-129 AF240786.1	11418522 NT	214585	214685	214585	1 0E-129 AB040892 1		1.0E-129 AW755254.1	1.0E-129 AW 765254.1	
Most Similar (Top) Hit BLAST E Value	1.0E-128		1.0E-128	1.0E-128	1.0E-128	1.0E-128		ì		1.0E-128 /		1.0E-128	1 0E-128	1 0F-128	1.0E-128	1.0E-128	1.0E-129 S37722.1	1.0E-129	1.0E-129		1.0E-129	1.0E-1291	1.0E-129	1.0E-129 Q14585	1.0E-129 Q14585	1 0E-129 Q14585	1 OF-120			l	١
Expression Signal	6.14	0.7	0.85	0.85	2.58	6.9	0.62	79.0	0.73	0.73	3.1	3.62	3.24	136	4.	7.02	1.33	1.19	2.73		1.57	1.67	2.78	121	121				2.26		
ORF SEQ ID NO:	29988	31147					L	33471		34277	35903	l	}				25839	١			27158	27169						1	29611		
SEQ ID NO:	17361	18257	18672	18672	19094	19493	20010	20356	21138	21138	22685	23302	ı	11505	23446	24840	L	L	L		14460	14460			1_	L	1	888	16988		ı
Probe SEQ ID NO:	4848	5458	9889	9889	6324	6831	7327	7692	8446	8446	10037	1080R	2007	01001	10093	24.5	418	404	1713		1717	4747	1838	34.25	34.28	27.5	3120	4143	4247	4247	127

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Prodect Earl 1977 1979 1970			,	_		_	_	_	_		_	Т	7	Т	Т	т		Τ	ΤŤ	Ť	Ť	T	Ť	Ť	Ť	Ĩ	Ĩ		T	Ī	T	T	٦	•	
Exam Noct Signal Most Similar Top Hit Acesslom Top		Top Hit Descriptor	Homo sapiens KVLQT1 gene	601449740F1 NIH_MGC_65 Homo sapiens CDINA clone IMAGE:3853688 5	601449740F1 NIH MGC 65 Homo septens dulta ciale in 1	Homo sapiens KVLQ11 gene	Homo saplens similar to ribosomal protein 320 (The s	Homo sepiens W SCK4 gene, excits 5 and 4	Homo saplens workey gate, however the same and the same same and same same same same same same same same	Homo sapiens miXNA for NIAANOS4 process, process, process, member 9 (SLC21A9), mRNA	Homo sapiens some family 21 (organic anion transporter), member 9 (SLCZ/NS), minor	Homo saplens solute carrier rating 2.1. Homo saplens cDNA clone IMAGE:1020288 3	ae91c01.31 Orangadre con Anno Sapiens CDNA clone IMAGE:1047589 5	af7207.r.1 Source, Villament american Sp6 (H. saplens) (LOC63694), mRNA	Home sapiens similar to ribosonna process of 79AA 1001410 5'	AU143115 YEAA I DUILU SEPERATORIA CIONE Y78AA1001410 5'	AU143115 Y79AA1 Homo sapiens Corrections and August Constitution of the Sapiens Consti	y49005.1 Soares feet all the Specific Annual Annual ASIATICUS=ASIAN	SP:B48150 B48150 HF-28=hBEAVA	Homo saplens hypothetical protein (HSP CA44), illings	Homo saplens mRNA for KJAA1414 protein, parter 3346366 5	601121995F1 NIH MCC_ZO Trans and A clane IMAGE:3346366 5	601121995F1 NIH MIGC 20 Halling Septemble to chromosome 11, band p13	Human gene for catalase (CO : : : : : :	Hambischer All Mac 53 Homo septens cDNA clone IMAGE:3685466 5	601343010F1 NIH MGC_53 Home sapiens cDNA clone IMAGE:3685466 5	Homo sapiens retinol dehydrogenase homolog isotorm-1 (NO.1) IIII 4.1.1	Т	Г						
Expn ORF SEQ Expression Most Similar Implied No. No. No. SEQ ID ID NO: Signal IPAST E ID HIT Aces Signal IDAST E ID HIT Aces No. No. No. No. No. No. No. No. No. No.	TION A	Top Hit Database Source		T HUMAN	Г			L	LN LN	NT	NT	ı	EST_HUMAN	EST HUMAN	ΝΤ	EST HUMAN	EST_HUMAN		EST HUMAN	. 1	NT	EST HUMAN	EST HUMAN	N	TN	EST HUMAN	TONIOL TOU	EST HUMAN	EST HUMAN	EST HUMAN	N.	EST HUMAN	EST HUMAN	EST_HUMAN	
Exam ORF SEQ Expression (Top) SEQ ID ID NO: Signal (Top) NO: 19781 3.73 1.06 19477 32499 0.56 1.00 19477 32499 0.56 1.00 19779 32191 2.49 1.0 20105 33191 2.49 1.0 20105 33191 2.49 1.0 20105 33191 2.49 1.0 20105 33192 2.49 1.0 20105 33191 2.49 1.0 20105 33191 2.49 1.0 20105 33192 2.49 1.0 20106 33192 2.49 1.0 20107 32844 6.57 1.1 30107 32836 0.97 1.0 30107 32842 0.57 1.1 30107 32842 0.57 1.1 4017 22823 1.38	Buis	op Hit Acession No.	4 2000	J000343.1	T		44420850	F041056.1	Γ		11437282	11437282	AA682200.1	A AR25528 1	11420850	A11443115.1	A1143115.1			AL 120/3	AR03783		BE275192.1	X04092.1	AJ010230.1	DE564219.1) BE564219.1	0 AF240698.1	0 BE 304219.1	0 BE3042 (9. 1	0 0407740 1	D AW843993.1	MAW363299.1	AW363299.1	2000
Expn ORF SEQ Expression Month SEQ ID ID NO: Signal B NO: 19781 3.73 19477 32499 0.56 19477 32499 0.56 19779 32500 0.56 19779 3276 4.15 20105 33191 2.49 20105 33191 2.49 20105 33191 2.49 20105 33192 2.49 20105 33192 2.49 20105 33192 2.49 20105 33192 2.49 20105 33192 2.49 20105 33192 2.49 20106 33193 0.57 20107 32836 0.57 32844 6.57 1.38 4 24213 37536 1.38 5 24213 37538 1.38 6 24213 37536 1.38 74<		st Similar op) Hit AST E	-	1.0E-129 A	1.05-129	u L	-			_	-	1 OF-129	_	_		00.1	1.05-128	1.05-120	1.0E-129	1.0E-129	1.0E-130	1.05-130	1.00-130	-10E-130	1 OE-130	1.0E-130	1.0E-13(1.0E-13	1.0E-13	1.05-13	1			1	╛
E wn ORF SEQ DOO: 19781 31742 19719 32498 19719 32498 19719 32498 19719 32700 33191 32835 33192 22627 33835 33192 22627 33835 33192 22627 33835 33192 22627 33835 19719 32844 19779 32844 17801 19740 22659 197407 27098 197407 297407 297407 297407 207407 197407 27098				3.73	0.56	0.56	- 1	1.	2.48	2.49	100	70.0	0.97	VC.0	4 4	0.57	1.38	1.38	1.79	2.68					3.00										
SEO ID SE				31742	32499	32500	32776	32844	33191	33192		35835	35836	36396	37137	32844										1	١								
				18781	19477	19477	19719	19779	20105	1 1		1)		L	Ш		1_	1	1	1	1_	L		\Box		. L	1	\perp	L		Ι.	<u></u>	L	L	l
			ö	9000	8846	6816	7027	7080	7428	7428	8217	8979	9979	10523	11186	11269	11615	11615	12106	1248		114	186	166	197	277	8 8	8 1 2	1,5	3 18	39	4	\$	18	188

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Single Exol Flobes Explessed ii Brail	Top Hit Descriptor	Human germline immunoglobulin lambda light chain pseudogene (VII.1)	CMO-CN0045-170200-225-g03 CN0045 Homo seplens cDNA	CM0-CN0045-170200-225-303 CN0045 Homo sepiens cDNA	Homo sapiens estrogen-responsive B box protein (EBBP), mRNA	Homo sapiens solute carrier family 6 (neurotransmitter transporter, L-proline), member 7 (SLC6A7), mRNA	Homo sapiens aurora-related kinase 1 (ARK1) mRNA, complete cds	EST368312 MAGE resequences, MAGD Homo saplens cDNA	Homo sapiens mRNA for KIAA1335 protein, partial cds	xd36e06.x1 NCI_CGAP_Ov23 Homo sapiens cDNA clone IMAGE:2595874 3'	Homo saplens contactin 6 (CNTN6), mRNA	Homo sapiens contactin 6 (CNTN8), mRNA	Homo sapiens hypothetical protein FLJ20208 (FLJ20208), mRNA	Homo sapiens hypothetical protein FLJ20208 (FLJ20208), mRNA	Homo sapiens glutamate receptor, metabotropic 5 (GRM5) mRNA	Homo saplens RET finger protein-like 1 antisense transcript, partial	과38c04.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:667590 당 similar to TR:G222811 G222811 ALPHA 1 CHAIN OF TYPE XII COLLAGEN ;	2758c04.r1 Soares_NHHMPu_S1 Home saplens cDNA clone IMAGE:667590 5' similar to TR:G222811 G222811 ALPHA 1 CHAIN OF TYPE XII COLLAGEN .	Homo saplens checkpoint suppressor 1 (CHES1), mRNA	Homo sapiens hypothetical probain FLJ20371 (FLJ20371), mRNA	Homo seplens hypothetical protein FLJ20371 (FLJ20371), mRNA	Homo sapiens DCRR1 mRNA, partial cds	Homo sapiens DCRR1 mRNA, partial cds	Homo sapiens beta-tubulin mRNA, complete cds	Homo saplens Cdc42 effector protein 2 (CEP2), mRNA	Human heparin cofactor II (HCF2) gene, exons 1 through 5	Homo saplens RNA-binding protein S1, serine-rich domain (RNPS1), mRNA	Homo saplens mRNA for multidrug resistance protein 3 (ABCC3)	Homo saplens mRNA for multidrug resistance protein 3 (ABCC3)	HUM516H08B Human placenta polyA+ (TFujiwara) Homo sapiens cDNA clone GEN-616H08 5'	HUM516H08B Human placenta polyA+ (TFujiwara) Homo sapiens cDNA clone GEN-516H08 5'	Human ribosomal protein L7 (RPL7) mRNA, complete cds
OIL LIOY BIE	Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	TN	TN	۲N	EST_HUMAN	IN	EST_HUMAN	TN	TN	ΙN	IN	TN	NT	EST HUMAN	FST HUMAN	N	N	TN	LN	TN	NT	IN	LΝ	NT	ΤN	NT	EST_HUMAN	EST HUMAN	LZ.
illo I	Top Hit Acession No.	30 X57825.1	30 AW843875.1	1.0E-130 AW843875.1	11425446 NT	11416777 NT	30 AF008551.1	30 AW956242.1	1.0E-130 AB037756.1	1.0E-130 AW103464.1	11432889 NT	11432889 NT	8923197 NT	8923197 NT	4504142 NT	1.0E-130 AJ010230.1	00 AA228126.1	0 0F+00 44228126 1	4885136 NT	8923349 NT	8923349 NT			0.0E+00 AF141349.1	32997		6857825 NT					00 L16558.1
	Most Similar (Top) Hit BLAST E Value	1.0E-130 >	1.0E-130/	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	0.0E+00	00=+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D83327.1	0.0E+00 D83327.1	0.0E+00	0.0E+00	0.0E+00 M58600.1	0.0E+00	0.0E+00 Y17151.2	0.0E+00 Y17151.2	0.0E+00 D78804.1	0.0E+00	0.0E+00L
	Expression Signal	0.55	0.81	0.81	0.75	2.62	0.45	2.39	1.64	1.25	0.51	0.51	1.72	1.72	2.67	1.56	9:	9	1.02	0.72	0.72	6.7	6.7	17.04	1.19	0.82	4.22	0.76	0.76	3.04	3.04	5.76
	ORF SEQ ID NO:	32405	32587	32588		32910		34553	34967				37194	37195	37624		26445	<u> </u>		26454	25465			25468	25478	25481	25486		25513		26615	
	Exon SEQ ID NO:	19391	19557	19557	19570	19841	21274	21410	21802	22484	23109	23109	23904	23904	24298	15478	12832	12832	12834	12841	12841	12848	12848	12853	12861	12863	12867	12884	12884	12885	12885	12886
	Probe SEQ ID NO:	6299	6723	6723	6736	7154	8582	8718	9114	9833	10463	10463	11242	11242	11703	12769	4			44	14	8	20	25	33	36	စ္တ	99	33	99	99	24

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. Top Hit Descriptor	or48e07.x1 Jia bone marrow stroma Homo sapiens cDNA clone HBMSC_cr48e07.31	α 48e07.x1 Jia bone maπow stroma Homo saplens cDNA clone HBMSC _cr48e07.31	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Homo saplens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sepiens protein tyroslne phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sapiens amiloride binding protein 1 (amine oxidase (copper-containing)) (ABP1), nuclear gene	encoding mitochondrial protein, mRNA	Homo saciens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	Homo sapiens actin, beta (ACTB) mRNA	Human polyhomeotic 1 homotog (HPH1) mRNA, partial cds	HA1347 Human fetal liver cDNA library Homo saplens cDNA	Homo saplens mRNA for KIAA1363 protein, partial cds	H.sapiens noct gene (exon 2)	Is38b05.x1 NCI_CGAP_UM Homo sapiens cDNA clone IMAGE:2230833 3' similar to TR:Q99651 Q99651 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;	1638b05.x1 NCI_CGAP_UM Homo saplens cDNA clone IMAGE:2230833.3' similar to TR:099651 099651 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;	yy01h09.r1 Soares melanocyte 2NbHM Homo saplens cDNA clone IMAGE:270017 5'	yy01h09.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone iMAGE:270017 5'	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	A		ya83g04,r2 Stratagene fetal spleen (#937205) Homo saplens cDNA clone IMAGE:68310 5'	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	601460375F1 NIH_MGC_66 Homo sapiens cDNA done IMAGE:3863803 5'	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	d unknown genes			3E:345201 5' strailar to	
Top Hit Database Source	EST_HUMAN	EST_HUMAN	TN.	LN	TN	TN	ΝΤ	TN		NT	LN	TN	IN	EST_HUMAN	NT	NT	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	TN	NT	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	N T	F	EST_HUMAN	EST_HUMAN		
Top Hit Acession No.	AW069534.1	AW069534.1	M60676.1	M60676.1	4758977 NT	4758977 NT	4758977 NT	4758977 NT		4501850 NT	450444 NT	5016088 NT	U89277.1	Al114743.1	AB037784.1	X91213.1	AI623701.1	AI623701.1	N36040.1	N36040.1	4505938 NT	4505938 NT	T56945.1	T56945.1	450444 NT	BF036881.1	450444 NT	AF111168.2	BE295973.1	BE295973.1	W73973.1	
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	00+30.0	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	-	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	
Expression Signal	12.55	12.55	1.5	0.91	10.36	10.36	10.18	10.18		0.78	15.25	17.68	23.26	3.51	1.72	1.33	0.89	1.47	1.48	1.48	4.38	4.38	1.29	1.29	8.88	2.1	25.83	1	1.15	0.88	5.4	
ORF SEQ ID NO:	25519	25520	25524				25536	25537		25544				25562			25572	25572	25573	25574		25587	25593	72594		25609		25612	25613	25613	Ιİ	
Exan SEQ ID NO:	12888	12888	12891	12893	12900	12900	12900	12900		12906	12907	12916	12919	12925	12926	12928	12935	12935	15513	15513	12943	12943	12950	12950	12964	12968	12970	12973	12975	12975	12976	
Probe SEQ ID NO:	59	29	ន	65	73	73	78	9/		8	81	06	8	88	100	105	113	114	115	115	128	128	136	136	149	153	155	158	160	161	162	

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Single Exon Probes Expressed in Brain

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Top Hit Descriptor	DV2-HT0457-140200 089 404 HT2457-1	AV31110437-1142200-008-004 F110457 Home sapiens cDNA	Home scalars size 6	Home control stand inger protein mKNA, complete cds	Homo septens chromosome 21 segment HS21C002
 | Homo soniene mPNA for VIA Actor | Homo saplens mBNA for KIA Agged - 447. | Hillings dammes challenges out 1.4.07.000. | Homo septens CTC Hims and a control of a con | Homo sanians CTC + tract and an act of the sanians CTC + tract and the sanians CTC + t | Jomo carlena chromosom V 101 0 0
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	Top Hit Descriptor	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo saplens hypothetical protein (LOC51250), mRNA	Homo sapiens DCRR1 mRNA, partial cds	Homo sapiens DCRR1 mRNA, partial cds	Homo sapiens DCRR1 mRNA, partial cds	IL2-CT0031-181199-020-B03 CT0031 Homo sapiens cDNA	Homo sapiens potassium inwardly-rectifying channel, subfamily J, member 15 (KCNJ15) mRNA	Homo saplens potassium inwardly-rectifying channel, subfamily J, member 15 (KCNJ15) mRNA	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo sapiens ribosomal protein S5 (RPS5) mRNA	zv18c06.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:753994 5'	Homo sapiens SON DNA binding protein (SON) mRNA	Homo sapiens SON DNA binding protein (SON) mRNA	Homo saplens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens hormonally upregulated neu fumor-associated kinase (HUNK), mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens myeloid/lymphoid or mbæd-lineage leukemla (trithorax (Drosophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo sapiens X-box binding protein 1 (XBP1) mRNA	Human zinc finger protein zfp31 (ਟਾਂ31) mRNA, partial cds	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Human mRNA for KIAA0184 gene, partial cds	Human mRNA for KIAA0184 gene, partial cds	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	1000899 5'	Homo saplens mRNA for KIAA1019 protein, partial cds	AGE:2018457 3' similar to gb:X54199	RC2-CT0320-300100-016-a09 CT0320 Homo sapiens cDNA
Top Hit	Database Source	TN	LN.	L	۲	۲	EST_HUMAN	LN	NT	NT	LN	TN	EST_HUMAN	NT	NT	NT	LN	LN	LΝ	NT	NT	LN	TN	LN	TN	LN	LN.	N	Ŋ	EST_HUMAN	IN	EST_HUMAN	EST_HUMAN
Top Hit Acession	Ö	4507500 NT	7706028 NT	D83327.1	D83327.1	D83327.1	AW845293.1	4557029 NT	4557029 NT	AB028942.1	AB028942.1	4506728 NT	AA480002.1	4507152 NT	4507152 NT	AF114488.1	7657213 NT	7657213 NT	5174574 NT	4827057 NT	U71600.1	AF231919.1	AF231919.1	AF231919.1	4507500 NT	4503854 NT	D80006.1	D80006.1	4507500 NT	AU134963.1	AB028942.1	Al363014.1	AW754180.1
Most Similar (Top) Hit	BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00
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Hamo sapiens phosphoribosylglycinamide formyltransferase, phosphoribosylglycinamide synthetase, H.sapiens gene for RNA pol II largest subunit, exons 23-29 Homo sapiens ribosomal protein L19 (RPL19) mRNA yg09a02.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:31652 AU132898 NT2RP4 Homo sapiens cDNA clone NT2RP4000837 6' 601274981F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3816789 6' 601111520F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3352348 5 Homo sapiens 5-hydroxytryptamine (serotonin) receptor 18 (HTR1B) mRNA Homo sapiens 5-hydroxytryptamine (serotonin) receptor 18 (HTR1B) mRNA Homo sapiens keratin 18 (KRT18) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Mus musculus truncated SON protein (Son) mRNA, complete cds Top Hit Descriptor Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA Homo sapiens Interferon gamma receptor 1 (IFNGR1) mRNA H.saplens gene for RNA pol II largest subunit, exons 23-29 H.saplens gene for RNA pol II largest subunit, exons 23-29 H.sapiens gene for RNA pol II largest subunit, exons 23-29 phosphoribosylaminolmidazole synthetase (GART) mRNA Homo septiens ribosomal protein SS (RPS5) mRNA Homo septiens mRNA for KIAA1019 protein, partial ods Homo septiens SON DNA binding protein (SON) mRNA Homo septiens SON DNA binding protein (SON) mRNA Homo sapiens chromosome 21 segment HS21 C046 Homo sapiens mRNA for KIAA 1209 protein, partial cds EST27054 Cerebellum If Homo saplens cDNA 5' end Homo sapiens chromosome 21 segment HS21C046 Homo sapiens chromosome 21 segment HS21C046 Homo sapiens keratin 18 (KRT18) mRNA EST_HUMAN EST_HUMAN EST_HUMAN HUMAN HUMAN Top Hit Database Source EST 4503680 NT 4503680 NT 4503680 NT 4503914 NT 눌 4557879 NT 4504532 NT ż 뉟 4506728 NT 4507152 NT 4504532 N 4503680 N 4503680 N 4503680 N 4507152 4503680 N 4557887 4506608 Top Hit Acession 0.0E+00 AL163246.2 0.0E+00 AU132898.1 AB028942.1 AA324262.1 BE264447.1 BE385144.1 0.0E+00 AF193607.1 0.0E+00 45 ġ 0.0E+00 X74870.1 0.0E+00 X74870.1 0.0E+00 R17795.1 X74870.1 0.0E+00 X74870. 0.0E+00 E 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 > 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0F+00 Most Similar (Top) Hit BLASTE Value 3.15 3.18 2.66 1.39 6.45 4.02 1.99 1.23 2.64 9.28 0.84 28.13 5.45 1.97 2.49 1.39 1.35 1.07 <u>ڊ</u> 1.58 6. Expression Signal 25888 25889 25895 25896 25897 25853 25854 25855 25865 26816 26817 25821 26823 25824 25825 25824 ORF SEQ ID NO: 25819 25820 25435 25851 13246 13286 13278 13172 13173 13173 13205 13209 13219 13257 13175 13178 13208 13241 SEQ ID 13175 13176 13178 13178 13182 13241 13174 12822 13206 13207 ë \$ \$ \$ 455 461 477 472 ₹ 8 491 461 Probe SEQ ID 386 387 387 389 389 389 397 392 393 393 393

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Single Exon Probes Expressed in Drain	Top Hit Descriptor	PM0-DT0065-130400-002-c06 DT0065 Homo sapiens cDNA	Novel human gane mapping to chomosome 1	Homo sapiens PC326 protein (PC326), mRNA	11.2-FT0159-070805-120-F07 FT0159 Homo sapiens cDNA	Homo sapiens chromosome 21 segment HS21C010	OV2-BT0635-160400-142-h05 BT0635 Homo sapiens cDNA	601764858F1 NIH MGC 63 Homo sapiens cDNA clone IMAGE:3996998 5	Homo sapiens mRNA for KIAA1476 protein, partial cds	Homo sapiens transcription elongation factor B (SIII), polypeptide 1-like (1 CEBTL) mrvvA	Homo saplens guanine nucleotide binding protein (G protein), alpha 11 (Gq class) (GNA11) mrvix	Homo sapiens guanine nucleotide binding protein (G protein), alpha 11 (Gq class) (GNA11) minner	Lamo caniana aniiin (LOC64443), mRNA	Homo saplens anillin (LOC54443), mRNA	Homo seniens X-linked anhidrottic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	THE BIT AND IT IS NOT COMP. Sub3 Homo saptens cDNA clone IMAGE:2713951 3'	Homo saniens RGH1 gene, retrovirus-like element	Home carriers inhining cytochrome c reductase. Rieske iron-sulfur polypeptide 1 (UQCRFS1), nudear gene	numb septent septent in RNA encoding introduction A-I (ApoA-I) gene, exon 1 Human apolipoprotein A-I (ApoA-I) gene, exon 1	R01822627F1 NIH MGC 75 Homo saplens cDNA clone IMAGE:4045447 6	Homo saciens hypothetical protein FLJ20701 (FLJ20701), mRNA	Homo saplens hypothetical protein FLJ20701 (FLJ20701), mRNA	Homo sapiens acetyl-Coenzyme A carboxylase beta (ACACB), mRNA	Homo saplens Smad- and Olf-interacting zinc finger protein mRNA, partial cas	Homo saplens Smad- and Olf-interacting zinc finger protein mRNA, partial cds	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mKNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mKNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mKNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mKNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mKNA	250007.r Soares tastis NHT Homo sapiens cDNA clone IMAGE: 726732.0	Home saplens RGH2 gene, retrovirus-like element	
e Exon Probe	Top Hit Database Source	HIMAN	Т		HIMAN	ולא <u>ל</u>	NAM.	NOWAL TOWARD	FO TOWNER								HUMAN	Z	TN	NAME TO T	HOLL TOWN	<u> </u>	12	FIN	2 12	Į.	Į.	IN IN	Į.	LV	LN	EST HUMAN	L	
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	Most Similar (Top) Hit BLAST E	- 18		0.0E+00 A	0.0=+00	0.0E+00 B	0.0E+00 A			0.0=+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00 D10083.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0=+00	0.0E+00	-	١	0.0=+00	0.05+00	0.05+00	0.0=+00			0.00
	Expression Signal		1:11	1.33	1.27	0.77	5.37	1.29	1.14	1.68	27.68	4.33	4.33	76.0	0.97	5.82	1.89	8	2.63	7	1.83	0.98	0.98	0.78	1.15	1.15	3.18	1.2	3,83	3.83				10.1
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Table 4 Single Exon Probes Expressed in Brain	. Top Hit Descriptor	2h61b04.r1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sepiens cDNA clone IMAGE:415567 S' similar to gb:A21187 ALPHA-2-MACROGLOBULIN PRECURSOR (HI MAN).	zh51b04.r1 Soares, fetal liver_spieen_1NFLS_S1 Homo saplens cDNA clone IMAGE:415567 6' similar to	Homo sablens novel SH2-containing models 3 Ais B3 - BNA	Homo sapiens diutemente recentor involtante in materialia.	Homo sapiens CCAAT-box-binding transcribition factor (CBE2) mBNA	Human neutral amino acid transporter (ASCT1) gene, exon 8	Homo sapiens sodium/calclum exchanger isoform NaCa3 (NCX1) mRNA, complete cda	Homo saplens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds	Homo saplens protein kinase, X-linked (PRKX) mRNA	riomo sapiens protein kinase, X-linked (PRKX) mRNA Himman andro	Home serious Hick mobility (FERVS)	Homo saplens mBNA for kit A4000	Homo sapians similar to refinitional moments and the sapians similar to refinitional moments.	np49d01.s1 NCI_CGAP_Br1.1 Home septens QDNA done IMAGE:1129633 3' similar to gb:X67362	Human von Willebrand factor cano 22 th	Human von Willebrand feder gene and 22 uituugil 34	Homo sapiens TNF recentor associated forces of 7 TP A E4 TP A E4	Homo saplens ALR-like protein mRNA harfiel case	Homo saplens ALR-like protein mRNA partial ods	Homo saplens hypothetical protein FLJ21634 (FLJ21634) mRNA	TCAAP1D0779 Pediatric acute myelogenous leukemia cell (FAB M1) Baylor-HGSC project=TCAA Homo	Appension of the contract of t	Houro septens WHC class I antigen (HLA-G) mRNA, HLA-G1 allele, complete cds	Homo sapians chloride channel O (4 (2)2). This	Human, plasmingren artifizator inhibitions, and a constant of the constant of	Human, plasminoden activator Inhibitor 1 gene, axons 2 to 9	Homo sapiens zinc finder protein 212 (ZNE212)	Homo saplens mRNA for repressor protein partiel rate	601445647F1 NIH_MGC_65 Home saplens cDNA clone IMAGE:3849803 6
ale Exon Pro	Top Hit Database Source	EST_HUMAN	EST HIMAN		LN L	LN.	NT	L	ĘŲ!	Z	Z	-	Į.	-	EST HIMAN	L	Į.		LN LN	Ę		HOT HIMAN	T							T_HUMAN
Sin	Top Hit Acession No.	8811.1	8811.1	4885526	F009009	5031624 NT	235.1	08389.1	08389.1	4820947	147.1	4504424	0.0E+00 AB029012.1	7468	0.0E+00 AA814537 1	ľ		32192	14750.1	4750.1	11545800 NT	1577.1	T	Ī		64.1	4.4	6912749 NT	30612.1 NT	П
	Most Similar (Top) Hit BLAST E Value	0.0E+00 W7	0.0E+00 W7	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U05	0.0E+00 AF1	0.0E+00 AF1	000	0.0E+00 X57	0.0E+00	0.0E+00	0.0E+00	0.0E+00.A	0.0E+00 M60675.1	0.0E+00 M60675.1	0.0E+00	0.0E+00 A	0.0E+00 AF26	0.0E+00	0.0E+00 BE24	0.0E+00 AF226990 2	0.0E+00 AF226990.2	0.0E+00 AF17	0.0E+00 J037	0.0E+00 J037	0.0E+00	0.0E+00 D30612.1	0.0E+00 BE869735.1
	Expression Signal	0.76	0.76	4.99	2.88	1.17	2.53	1.07	1.07	4 98	1.15	10.4	4.49	2.43	9.24	4.34	4.34	1.71	4.95	4.95	12.29	2.12	1.07	1.07	2.41	1.07	1.07	1.38	1.86	3.01
	ORF SEQ ID NO:	28041	28042		26054	28057	28061	20004	26070	26071		26084	26088	26101	26115	26119	26120	26129	26135	26136	26139	26146	26170	26171	26172	26175	26176	26177	26179	26180
	Exon SEQ ID NO:	13407	13407			13419	13422	13420	13431	13431	15547	13443	13448	13456	13468	13471	13471	13481	13486	3486	13488	13483	13512	13512	13513	13516	13516	13519	18851	13521
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Top Hit Descriptor y69g08.r1 Soares breast 2NbHBst Homo sapiers cDNA clone IMAGE:164046 5'	Homo sepiens splicing factor 3a, subunit 1, 120kU (3F3A1), mkwa Homo sepiens gene for AF-6, complete cds	Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	Human mRNA for KIAA0184 gene, partial cds	Human mRNA for KIAA0184 gene, partial cds	H.sapiens mRNA for interferon alpha/beta receptor (long form)	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens pericentrin (PCNT) mRNA	Homo sapiens T-cell lymphoma Invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens potassium voltage-gated channel, isk-related family, membar 1 (KCNE1) mRNA	Homo sapiens serine-threonine protein kinase (MNBH) mRNA, complete cds	Homo sapiens serine-threonine protein kinase (MNBH) mRNA, complete cds	Homo sapiens GA-bInding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Homo sapiens T-cell lymphoma Invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens T-oell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens sodium/myo-inositol cotransporter (SLC5A3) gene, complete cds	Homo saplens mRNA for KIAA1019 protein, partial cds	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo saplens SON DNA binding protein (SON) mRNA	Homo sapiens mRNA for KIAA1019 protein, partial cds	Homo sapiens ribosomal protein S5 (RPS5) mRNA	Homo saplens mRNA for KIAA0910 protein, partial cds	Homo sapiens mRNA for KIAA0910 protein, partial cds	nj66d07.s1 NCI_CGAP_Pr10 Homo saplens cDNA clone IMAGE:897453	nj66d07.s1 NCI_CGAP_Pr10 Homo saplens cDNA clone IMAGE:897453	602085579F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249915 5'	Homo saplens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo saplens hormonally upregulated neu fumor-associated kinase (HUNK), mRNA	Homo saplens hormonally upregulated neu tumor-essociated kinase (HUNK), mRNA
Top Hit Database Source EST_HUMAN		NT	NT	NT	NT	LN	LN	F	LN	١N	NT	LN	NT	NT	L	Ŋ	IN	LN	IN	LN	NT	M	NT	LΝ	ΙN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LN	N	F	F
Top Hit Acession No. R48915.1	5032086 NT AB011399.1 NT	7661965 NT	D80006.1	D80006.1	X89772.1	AB020717.1	AB020717.1	5174478 NT	4507500 NT	7857213 NT	7657213 NT	4557686 NT	AF108830.1	AF108830.1	. 4503854 NT	4507500 NT	4507500 NT	AF027153.1	AB028942.1	AB028942.1	4507152 NT	AB028942.1	4506728 NT	AB020717.1	AB020717.1	AA533272.1	AA533272.1	BF677694.1	7657213 NT	7657213 NT	7657213 NT	7657213 NT
	0.0E+00/				0.0E+00	0.0E+00/	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/			0.0E+00	0.0E+00		0.0E+00/	0.0E+00/	0.0E+00/		0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal 3.51	2.4	234	1.18	1.18	2.13	5.97	5.97	9.88	11.63	1.96	4.28	2.81	1.39	1.39	1.58	2.09	2.09	1.32	4.62	4.62	9.56	4.34	8.0	2.2	2.2	2.45	2.45	9.44	1.94	1.94	3.31	3.31
g _O □	26184	26196	26208	Ш				26222			26243			26252		26262			26274	26275			26278	26281	26282	26283			26285			
Exon SEQ ID NO: 13524	13525	13537	L		13552	13556	13556	, 13560	13561	13577	13578	13580	13585	13585	13590	13593	13593	13600	13604	13604	13605	13606	13607	13611	13611	13612	13612	<u> </u>	L	13617		13618
Probe SEQ ID NO: 752	38 28	\$	775	775	贸	784	18	788	789	805	806	808	814	814	819	823	823	88	834	834	835	836	837	841	841	842	842	쫎	847	847	848	8

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Homo saplens KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog (KIAA0929), mRNA Homo sapiens laminin receptor 1 (67kD, ribosomal protein SA) (LAMR1), mRNA Homo sapiens laminin receptor 1 (67kD, ribosomal protein SA) (LAMR1), mRNA Homo sapiens alpha-1-antchymotrypsin precursor, mRNA, partial cds Homo sapiens kallistatin (P14) gene, exons 1-4, complete cds Homo sapiens of cerdiac alpha-myosin heavy chain gene Homo sapiens of cerdiac alpha-myosin heavy chain gene Homo sapiens thyrotrophic embryonic factor (TEF), mRNA os99e03 s1 NCI_CGAP_GC3 Homo sapiens cDNA clone IMAGE:1613404 3' os99e03 s1 NCI_CGAP_GC3 Homo sapiens cDNA clone iMAGE:1613404 3' Homo saplens chromodomain protein, Y chromosome-like (CDYL) mRNA Humen beta-tubulin (TUB4q) gene, complete cds
Humen beta-tubulin (TUB4q) gene, complete cds
Humen beta-tubulin (TUB4q) gene, complete cds
Homo sapiens 8q22.1 region and MTG8 (CBFA2T1) gene, partial cds
Homo sapiens 8q22.1 region and MTG8 (CBFA2T1) gene, partial cds
Homo sapiens 14q32 JaggedZ gene, complete cds; and unknown gene Top Hit Descriptor QV0-BT0703-280400-211-911 BT0703 Homo sepiens cDNA QV0-BT0703-280400-211-911 BT0703 Homo saplens cDNA PM2-GN0014-050900-001-f02 GN0014 Homo saplens cDNA PM2-GN0014-050900-001-f02 GN0014 Homo sapiens cDNA PM2-GN0014-050900-001-f02 GN0014 Homo sapiens cDNA Homo sapiens thyrotrophic embryonio factor (TEF), mRNA Homo sapiens mRNA for KIAA0994 protein, partial cds Homo sapiens mRNA for KIAA0994 protein, partial cds Human ras inhibitor mRNA, 3' end Homo saplens chromosome 21 segment HS21C003 Homo sapiens chromosome 21 segment HS21C003 Human beta-tubulin (TUB4q) gene, complete cds Homo saplens mRNA for PSP24, complete cds Homo sapiens partial o-fgr gene, exons 2 and 3 Homo sapiens partial o-fgr gene, exons 2 and 3 Human ras inhibitor mRNA, 3' end Human ras inhibitor mRNA, 3' end Single Exon Probes Expressed in Brain EST HUMAN Top Hit Database EST_HUMAN EST_HUMAN EST_HUMAN EST HUMAN EST_HUMAN HUMAN 4504958 NT 7657266 NT ĽΝ Þ 뒫뉟 눌 Top Hit Acession 4504958 4507430 4507430 0.0E+00 AL163203.2 0.0E+00 BF368974.1 0.0E+00 BF368974.1 0.0E+00 BF368974.1 0.0E+00 X62207.1 0.0E+00 X52207.1 0.0E+00 AB023211.1 0.0E+00 AB023211.1 ġ 0.0E+00 BE089592.1 0.0E+00 AI001948.1 0.0E+00 AI001948.1 0.0E+00 AB030566. 0.0E+00 AF111170.3 0.0E+00 AF198490.1 0.0E+00 AF198490.1 0.0E+00 M37190.1 1,10187 Z20656.1 M37190.1 0.0E+00 M37190 0.0E+00 U83668.1 0.0E+00 U83668.1 0.0E+00 U83668.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 Vost Similar (Top) HII BLASTE ·0.0E+00 0.0E+00 0.0E+00 1.78 2.98 1.05 Expression 9.95 2.18 2.18 3.38 9.49 1.11 1.69 1.69 50.9 5.52 8.64 Signal 26316 26317 26326 ORF SEQ 26333 26337 26337 26355 26362 26363 28384 28385 ΘŅ 26375 26393 26392 26394 26395 26402 26410 26416 Exon SEQ ID 13640 13672 13709 ğ 15556 13727 13728 13748 13728 13749 13752 13727 13737 Probe SEQ ID 878 888 898 900 901 962 2 2 2 2 2 3 942 972 98 88 88 984 686 990 88

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	Top Hit Descriptor	Homo sapiens 14q32 Jaggedz gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds, and unknown gene	Homo sapiens 14q32 Jaggad2 gene, complete cds; and unknown gene	Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M0122), III W. Homo sapiens DKFZP586M0122 protein (DKFZF380M01222), III W. Homo sapiens DKFZF380M0122 protein (DKFZF380M01222), III W. Homo sapiens DKF	Homo sapiens Inner membrane protein, mitochondria (mitolinia) (with the sapiens Inner membrane protein, mitochondria (mitolinia) (with the sapiens Inner membrane protein, mitochondria (mitolinia) (with the sapiens inner membrane protein, mitochondria (mitolinia) (with the sapiens inner membrane protein, mitochondria (mitolinia) (with the sapiens inner membrane protein, mitochondria (mitolinia) (with the sapiens inner membrane protein) (mitolinia) (with the sapiens inner membrane protein) (mitolinia) (mito	aa88g07.s1 Stratagene fetal retina 937202 Homo sapiens CONN Giulo III.	SW.PRS8_HUMAN P47210 28S PROTEASE RECOUNTY OF THE PROPERTIES THE POLYMERASE II EST51124 WATM1 Homo septens CONA clone 51124 similar to DNA-DIRECTED RNA POLYMERASE II	(alignment Ser and Pro with BLASTX or P) (alignment Ser and Pro with BLASTX or P) EST51i24 WATM1 Homo saplens CDNA done 51i24 similar to DNA-DIRECTED RNA POLYMERASE II	(alignment Ser and Pro with BLAS ix or P) Home sapiens hypothetical protein FLJ11196 (FLJ11196), mRNA	Homo sapiens heat shock 70kD protein 9B (mortalin-2) (HSPA9B) mKNA	Homo sapiens cadherin 6, K-cacherin (fetal kidney) (CDRG) mixik	Homo saplens cadherin 6, K-cadherin (fetal kidney) (CDHs) mkNA	Homo sapiens hypothetical protein FLI20695 (FLI20695), mkNA	Homo sapiens hypothetical protein FLJ20695 (FLJ2099), minor	Homo saplens mRNA for alpha-tubulin d (1 Obro gene)	Homo saplens hypothetical protein PLJZUVOV (PLZZVVVV),	Homo sapiens alkylation repair; alko nomolog (Apr. 1),	Homo saplens Dearn associated promise Considerate Homo saplens cons	MRO-BN0115-20030C-003-100 BING 100 100 100 100 100 100 100 100 100 10	Homo sapiens pousseum creamers arthfamily K. member 9 (KCNK9), mRNA	Homo sapiens porassium ciratine, description (PRKX) mRNA	Home sapietis process and the second	Honio septemb presental profein S27a (RPS27A) mRNA	Homo sabiens hypothetical protein FLJ20309 (FLJ20309), mRNA	Homo saplens DNA for Human P2XM, complete cds	Homo saplens DNA for Human P2XM, complete cds	Homo sapiens similar to rat integral membrane grycoprocein BOM121 (BOM121 1). mRNA	Homo sapiens similar to rat integral membrane glycoprotein Fowner (1997)	Homo sapiens New38-binding protein typing (Co.	H.sapiens ART4 gene	
	Top Hit Database Source		Į.	LZ LZ	: -	<u> </u>		EST_HUMAN	EST_HUMAN	EST HUMAN	Z	2 2	Z	L'N	LZ	LN.	IN:	LN	ΙΝ	EST_HUMAN	TN1	4 NT	N.T	7N7	ZNZ	INO	Z LV	IN 8	N 88	NT OC	NT	
ignio.	op Hit Acession No.		T	T	THITTING	7861663 TM A116NAT	1000	AA458680.1	N43182.1	N43182.1	8922933 NI	VI 050001	48266/2 N	TN 482000	N 402360	4 1045020 4	TN TROCCOS	5174384 NT	4758117 NT	BE005208.1	7706134 NT	7706134 NT	4826947 NT	4826947 NT		8923290 N	0.0E+00 AB002059.1	0.0E+00 AB00203933		7706500 NT	0.0E-00 X95828.1	
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				۹L	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00 N	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.06+00	0.0=+00	0.00+000	0.00+00	00+100		_	0.0E+00	0.0E+00	00+30.0	0.0E+00			١	0.0E+00	1		
	Expression (Signal E		1.83	2.3	3.76	3.06	2.5	1.43	1.9	1.9	3.55	2.49	1.89	1.89	3.63	3.63	119.02	1.19	3.32	2.3	7.7						2			"		3 0.87
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	-	ÿ	13755	13755	13756	13759	l	i	13767	13787	1.	L.	L	\mathbf{L}_{-}	L	13806	13807						- 1	1	13867	1	L	1_	<u></u>	19 13876		23 13879
	Probe SEQ ID	ö	å	988	8	666	1003	1004	1007	100,	1010	1025	1043	1043	74	1047	1048	1050	1052	1060	1074	1097	1097	1110	113	1111	1116	1118	1119	1119	1122	1123

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Г	—т	Т	7	Т	Т	Т	Т	Т	Т	Т	Т	Т	Т	Т	Т	T	T	Т	Т	Ï	T	T	7	T	Ŧ	Ŧ	T	Ť	Ť	Ť	Ť	T	T	T	Ţ
	Top Hit Descriptor	H.saplens ART4 gene	db22d10.x1 Soeres_pregnant_uterus_NbHPU Homo eaptens cDNA clone IMAGE:169/011 3	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo sapiens chondroitin sulfate proteoglycan 2 (versican) (CSPG2) mRNA	Homo sapiens chondroltin sulfate proteoglycan 2 (versican) (CSPG2) mRNA	Homo sapiens chromosome 12 open reading frame 3 (C12ORF3), mRNA	Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD2s, mKNA	Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD23, miniva	Homo sapiens mRNA for KIAA1414 protein, partial cds	Homo saplens keratin 18 (KRT18) mRNA	Homo sapiens amphiphysin I mKNA, alternative splice isotorm, complete cos	Homo sapiens mutt. (E. coli) homolog 3 (MIH3), mKNA	Homo saplens hypothetical protein FLJ10697 (FLJ10897), mKNA	Homo sapiens ALR-like protein mRNA, partial cds	Homo saplens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo saplens ALR-like protein mRNA, partial cds	Homo saplens chromosome 3 subtelomeric region	Homo sapiens chondroitin sulfate proteoglycan 4 (metanoma-associated) (CSP'04), mrdNA	Homo saplens NF2 gene	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo sapiens Williams-Beuren syndrome deletion transcript 9 (WBSCR9) mKNA, compiete das	Homo sapiens mRNA for KIAA1507 protein, partial cds	Homo sapiens mRNA for KiAA1507 protein, partial cds	Homo sapiens Wolfram syndrome (W FS) mRNA	Homo sapiens Wolfram syndrome (W FS) mRNA	Homo sapiens Wolfram syndrome (WFS) mKNA	Homo sapiens protein phosphatase 24 Brt gamma subunit gene, exon 5	Homo saplens mabdoid tumor delican region protein region protein region protein region protein regions and region protein regions and regions are regions and regions and regions and regions and regions are regions and regions and regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions and regions are regions and regions are regions and regions are regions and regions are regions and regions are regions and regions are regions are regions and regions are regions are regions and regions are regions are regions and regions are regions are regions are regions are regions and regions are regions are regions are regions are regions are regions.	Homo sapiens rhabdold tumor deletion region protein 1 (RTDR1), mKNA	Homo sapiens ring finger protein 9 (KNF9), mKNA	Homo saplens zinc finger protein 173 (ZNF173) mKNA	Homo saplens ring finger protein 9 (RNF9), mKNA	Homo sapiens zinc ringer procein 173 (zunf 173) ilinuuh
	Top Hit Database Source	NT	EST HUMAN	Z	LZ	LN TN	LN LN	NT	ΤN	LN	NT	LN	NT.	TN	NT	۲N	IN	NT.	NT	LZ	F	F	LΣ	ΝΤ	NT	NT	۲	LN	LN	LN	LN	LN L	NT	LZ	LN
3	Top Hit Acession No:	X96826.1	-		4758081 NT	4758081 NT	9966844 NT	7305076 NT	7305076 NT	AB037835.1	4557887 NT	AF034996.1	7657338 NT	8922593 NT	AF264750.1		AF264750.1	AF264750.1	AF109718.1	4503098 NT	Y18000.1	4506718 NT	AF084479.1	AB040940.1	AB040940.1	5174748 NT	5174748 NT	5174748 NT	AF096156.1	7657529 NT	7657529 NT	5803146 NT	4508004 NT	5803146 NT	4508004 N
	Most Similar (Top) Hit BLAST E Value	0.0E+00				0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00.0			0.0E+00/			0.0E+00	0.0E+00		0.0E+00/		0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	0.87	000	1.56	0.77	0.71	1.39	6.83	6.83	1.85	0.75	4.54	1.52	1.7	1.53	1.63	177	1.63	9.12	171	233	23 69	3.07	2.07	2.07	2.04	2.04	2.04	3.78	1.63	1.63	2.03	0.89	1.12	0.72
	ORF SEQ ID NO:	26539	28840	28542	26548	26549	26550	26561	26562	26565	26574	26583		26617	26620							26667	L				26696	26697		26709					26718
	SEO ID NO:	13879	42000	13887	13880	13889	13890	13900	13900	13903	1		13939	13953	13956	13956	L	L	L		L	L	L		1	1	L	L		1_	ı	1		1	14046
İ	Probe SEQ ID NO:	1123		1124	133	1133	1134	1145	1145	1,48	1156	1167	1187	1201	1204	1204	1205	4200	1224	1000	4243	1243	125	1265	1285	1277	1277	1277	1278	1288	1288	1283	1204	1298	1297

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Single Exol Flobes Expressed in Digiti	Top Hit Descriptor	Homo sapiens mRNA for KIAA0577 protein, complete cds	Homo saplens KIAA0170 gene product (KIAA0170), mRNA	Homo saplens KIAA0170 gene product (KIAA0170), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Human endogenous retrovirus HERV-K10	Homo sapiens mRNA for Femilial Cylindromatosis cyld gene	Homo saplens partial TTN gene for titin	qg38b06.x1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:1837427.3' similar to WP:T27A1.5 CE14213;	RAN, member RAS oncogene family-homo sapiens RAN, member RAS oncogene family (RAN), mRNA	Homo saplens proprotein convertase subtilish/kextn type 2 (PCSK2) mRNA	Homo saplens proprotein convertase subtilisin/kexin type 2 (PCSK2) mRNA	Homo saplens KIAA1114 protein (KIAA1114), mRNA	Homo sapiens KIAA1114 protein (KIAA1114), mRNA	Homo sapiens partial AF-4 gene, exons 2 to 7 and Alu repeat elements	Homo saplens alpha1-6fuccsyltransferase (alpha1-6FucT) gene, exon 7	Homo sepiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Human nebulin mRNA, partial cds	Human nebulin mRNA, partial cds	Novel human gene on chromosome 20	Novel human gene mapping to chomosome 1	Human mRNA for KIAA0240 gene, partial cds	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo saplens KIAA0170 gene product (KIAA0170), mRNA	Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Human von Willebrand factor pseudogene corresponding to exons 23 through 34	Homo sapiens hHDC for homolog of Drosophila headcase (LOC51696), mRNA	EST371757 MAGE resequences, MAGF Homo sapiens cDNA	ae34e03.r1 NCI_CGAP_GCB1 Homo saplens cDNA done IMAGE:815116 5'	Cercopithecus aethiops cyclophilin A mRNA, complete cds
JIE EXUII PIO	Top Hit Database Source	NT	LN	LN	LN	NT	IN	TN	NT	EST_HUMAN	LΝ	LN	ĻΝ	LN	LΝ	TN	LN	NT	NT	LN	TN	NT	NT	NT	NT	NT	NT	NT	NŤ	NT	EST_HUMAN	EST_HUMAN	N L
	Top Hit Acession No.	AB011149.1	7661965 NT	7661965 NT	8567387 NT	8567387 NT	M14123.1	AJ250014.1	AJ277892.1	Al208756.1	6042206 NT	4505646 NT	4505646 NT	7705565 NT	7705565 NT	AJ238093.1	AF038280.1	4507720 NT	4507720 NT	U35637.1	U35637.1	AL 132999.1	AL137764.1	D87077.1	6912457 NT	7661965 NT	7661965 NT		M60676.1	7706434 NT			AF023860.1
	Most Similar (Top) Hit BLAST E Vatue	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00[0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00	0.0E+00						0.0E+00
	Expression Signal	4.71	2.04	5.88	3.01	3.01	1.82	1.38	3.39	1.59	13.21	2	2	4.08	4.08	4.59	3.02	5.39	5.39	17.93	17.93	2.59	1.82	1.73	4.53	1.55	1.55	0.97	0.97	1.37	1.21	1.76	49.82
	ORF SEQ ID NO:	26720	28721	26722	26723	26724	26737	26809	26817	26821	26822	26833	26834	26837	26838	26839	26852	26863	26864	26868	26869	26876	26877	26882	26885	26887	26888	26893	26894	26932	26949	26950	26953
	Exon SEQ ID NO:	14048	14049	14050	14051	14051	14062	14134	14140	14143	14144	14153	14153	14155	14155	14157	14168	14178	14178	14183	14183	14193	14194	14198	14201	14203	14203	14207	14207	14246	14263	14264	14269
	Probe SEQ ID NO:	1289	1300	1301	1302	1302	1314	1387	1393	1396	1397	1406	1406	1408	1408	1410	1420	1431	1431	1436	1436	1446	1447	1451	1454	1456	1456	1460	1460	1500	1516	1517	1522

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Table 4

Single Exon Probes Expressed in Brain	Top Hit Descriptor	Cercopithecus gethions cyclophilip 4 mBNA complete ada	EST388208 MAGE resembnes MAGN Lower contract PALA	EST388206 MAGE resemiences MAGN Homo september about	Bovine mRNA for neurocalcin	Homo saplens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomai protein (L441) and FTP3 (FTP3) canas complete add	Homo sapiens transmembrane divicionate in (GPNMR) mRNA	Homo sapiens transmembrane alycoprofein (GPNN/R) mBNA	Homo sapiens KIAA0967 protein (KIAA0967), mRNA	Homo saplens TNF-inducible protein CG12-1 (CG12-1) mRNA	Human transglutaminase mRNA. complete cds	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Homo saplens ribosomal protein L5 (RPL5) mRNA	Human laminin receptor (2H5 epitone) mRNA 5' end	Homo saplens titin (TTN) mRNA	Homo saplens titin (TTN) mRNA	Homo saplens chondroitin sulfate proteoglycan 4 (melanome-associated) (CSDO4)DNA	human o-yes-2 gene	H.sapiens hH2B/e gene	Homo sapiens butyrophilin, subfamily 2. member A1 (BTN2A1), mRNA	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	AV690831 GKC Hamo sapiens cDNA clane GKCBOF02 5'	AV690831 GKC Hamo sapiens cDNA clone GKCBOF02 5'	Homo saplens mRNA for KIAA1472 protein, partial cds	Homo saplens DNA polymerase zeta catalytic subunit (REV3) mRNA complete cds	Homo saplens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens KIAA0569 gene product (KIAA0569) mRNA	Homo sapiens heat shock 70kD protein 10 (HSC71) (HSPA1n) mRNA	Homo saplens heat shock 70kD protein 10 (HSC71) (HSPA1n) mRNA	Human sodium channel mRNA	yo76c05.s1 Soares adult brain N2b4HB56Y Homo sapiens cDNA clone IMAGE 183848 31	Homo sapiens mRNA for KIAA1609 protein, partial cds	Homo sapiens mRNA for KIAA1609 protein, partial cds
gle Exon Prob	Top Hit Database Source	L	T HUMAN	Т	Г	L					±N				- E				TN.	H				EST_HUMAN A								T_HUMAN	Г	
Sin	Top Hit Acession No.	+00 AF023860.1			-00 D10884.1	-00 U78027.1	4505404	4505404 NT	7662405 NT	7656972 NT		1507720	4507720 NT	4506654 NT		4507720 NT	4507720 NT	4503098 NT			5921460 NT	921460				0.0E+00 AF157476.1	7662183 NT	7662183 NT	5729876 NT	5729876 NT				П
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M98478.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D00333.1	0.0E+00 Z83738.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M91803.1	0.0E+00 H26973.1	0.0E+00 AB046829.1	0.0E+00 AB046829.1
	Expression Signal	49.82	1.24	1.24	5.49	2.07	2.1	2.1	3.3	7.29	1.84	3.72	3.72	11.72	11.72	2.86	2.86	11.7	1.21	1.65	1,59	1.59	11.18	11.18	9.85	1.83	9.78	9.78	42.75	42.76	7.94	5.85	2	2
	ORF SEQ ID NO:	26954	26957	26958	26959		26962	26963	28965		26971	26973	26974		26975	26988	26989	26990		27007	27008	27009	27010	27011	27014	27015	27018	27019	27020	27021	27023	27039	27051	27052
	Exan SEQ ID NO:		14271	14271	14272	14274		١	- 1	14278	14284	14287	14287	15572	14288	14300	14300	14302	14310	14321	14322	14322	14323	14323	139/3	14329	14331	14331	14333	14333	14335	14360	14381	14361
	Probe SEQ ID NO:	1522	1524	1524	1525	1527	1528	1528	1530	1531	1537	흅	1540	2	1542	1683	1563	1885	1583	1674	1575	1575	9/61	19/6	B/QL	1583	1586	1585	1587	1587	1589	1604	1614	1614

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Top Hit Descriptor	MRo-HT0168-191199-004-b11 HT0166 Homo sapiens cDNA	MR0-HT0166-191199-004-b11 HT0166 Homo sapiens cDNA	wg81b07.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2371477 3' similar to TR:062788 062788 CYS2HIS2 ZINC FINGER PROTEIN :	Homo saplens hematopoletic-derived zinc finger protein (HD-ZNF1) mRNA	Homo sapiens T-cell receptor gamma V1 gene region	Human zino-finger protain 7 (ZFP7) mRNA, complete cds	Human zinc-finger protein 7 (ZFP7) mRNA, complete cds	Homo sapkens keratin 18 (KRT18) mRNA	Homo saplens v-ets avian erythroblastosis virus E28 oncogene related (ERG), mRNA	hu11d05x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3168281 3' similer to TR:095147 095147 NRP-1 LIKE PROTEIN TYROSINE PHOSPHATASE;	hu11d05x1 NCI_CGAP_Lu24 Hamo saplens cDNA clone IMAGE:3166281 3' similar to TR:095147 095147 NKP-1 LIKE PROTEIN TYROSINE PHOSPHATASE;	Homo saplens gamma-aminobutyric acid (GABA) A receptor, gamma 2 (GABRG2) mRNA	yo59e08.r1 Soares breast 3NbHBst Homo septens cDNA clone IMAGE:182246 5' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN):	yo59e08.r1 Soares breast 3NbHBst Homo sapiens cDNA clone IMAGE:182246 6' similar to gb:M64099 GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (HUMAN):	H.saplens H2B/h gene	H.sapiens H2B/h gene	Homo sapiens high-mobility group (nonhistone chromosomal) protein 17 (HMG17), mRNA	Homo saplens WNT16 protein (WNT16) mRNA, complete cds	Homo saptens FOXJ2 forthead factor (LOC55810), mRNA	Homo saplens RNA binding motif protein, Y chromosome, family 1, member A1 (RBMY1A1) mRNA	Homo saplens WAVE2 mRNA for WASP-family protein, complete cds	TCR zeta [human, Genomic/mRNA, 365 nt, segment 1 of 8]	Homo saplens NOD2 protein (NOD2), mRNA	Homo saplens SMCY (SMCY) gene, complete cds	Homo saplens ribosomal protein S2 (RPS2) mRNA	Homo sapiens E1A binding protein p300 (EP300) mRNA	Homo saplens E1A binding protein p300 (EP300) mRNA	Human CSF-1 receptor (FMS) gene, complete cds, and (SMF) gene, partial cds	zd66g09.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:345664 5'
Top Hit Database Source	EST_HUMAN	EST_HUMAN	FST HUMAN	Į.	Ę	Ę	Ą	F	F	EST_HUMAN	EST HUMAN		EST HUMAN	EST HUMAN	N	LΝ		전	NT.	LN	LN	LN	TN	L	TN	N	T		EST_HUMAN
Top Hit Acession No.	BE144364.1	BE144364.1		58513	AF057177.1	M29580.1	M29580.1	4557887 NT	7657065 NT	BE222374.1	BE222374.1	57610	H30132.1			Z80780.1	5031748 NT	AF169963.1	8923841 NT	4826973 NT	AB026542.1	S94400.1	11545911 NT	AF273841.1	4506718 NT	4557556 NT	4557556 NT		W76571.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	00+00			0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.05+00		_		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00
Expression Signal	1.25	1.25	1.68	12	2.61	1.76	1.76	1.35	1.5	1.12	1.12	3.24	3.18	3.18	1.32	1.32	13.71	1.11	3.53	1.11	4.08	219	1.18	1.93	98'9	3.37	3.37	1.42	1.2
ORF SEQ ID NO:	27095	27096	27100	27101	27102	27106	27107	27109	27110	27113	27114	27116	27119	27120	27122	27123		27133	27136	27145	27162		27167	27181		27228	27229	27232	
SEQ ID	14405	14405	14409	14410	14411	14414	14414	14418	14417	14420	14420	14421	14424	14424	14428	14428	14429	14437	14438	14445	14451	14453	15577	14481	15578	14523	14523	14525	14528
Probe SEQ ID NO:	1659	1659	1663	1664	1665	1669	1669	1671	1672	1675	1675	1677	1680	1680	1682	1682	1685	1693	1695	1702	1708	1710	1725	1739	1778	1782	1782	1784	1787

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Single Exon Probes Expressed in brain	Top Hit Descriptor	Homo sapiens glutathlone S-transferase theta 2 (GS 1 /2) and glucannoing Commissions of the sapiens genes, complete cds	Human topolsomerase I pseudogene 1	Homo sapiens histidine ammonia-lyase (HAL) mRNA	Homo sapiens histidine ammonla-lyase (HAL) mRNA	Homo sapiens chromosome 21 segment HS21C052	Homo sapiens nebulin (NEB), mRNA	Homo sapiens nebulin (NEB), mRNA	Homo sapiens actinin, alpha 4 (ACTN4) mRNA	Homo sepiens actinin, alpha 4 (ACTN4) mRNA	Home sapiens mRNA for KIAA0790 protein, partial cds	Home sapiens mRNA for KIAA0790 protein, partial cds	Hirman TFEB protein mRNA, partial cds	TEFR nortein mRNA, partial cds	Jacob 1 NCI CGAP Pan1 Homo sapiens cDNA clone IMAGE:2679913 3'	Association CAP Part Hamp sapiens cDNA clone IMAGE:2879913 3'	Home seniors calcineurin binding protein 1 (KIAA0330), mRNA	Trans september in binding protein 1 (KIAA0330), mRNA	Houring september KIAA0408 gene product (KIAA0408), mRNA	Homo sapiens mRNA for KIAA0577 protein, complete cds	H sanians genes for semenogelin I and semanogelin II	H. sapiens genes for semenogelin I and semenogelin II	Homo sapiens mRNA for KIAA1513 protein, partial cds	Homo sapiens SMCY (SMCY) gene, complete cds	Homo sapiens SMCY (SMCY) gene, complete cds	Hamo saplens TP53TG3a (TP53TG3a), mRNA	AU140831 PLACE4 Homo saplens cDNA clone PLACE4000321 5	Homo sapiens KIAA1114 protein (KIAA1114), mRNA	Homo saplens KiAA1114 protein (KIAA1114), mRNA	7822E10 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7822E10	7R22E10 Chromosome 7 Fetal Brain cDNA Library Homo sapiens cDNA clone 7822E10	Homo saplens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo saplens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE6A), mRNA	HSC0IC021 normalized Infant brain cDNA Homo sapiens cDNA clone c-01c02	
e Exon Probe	Top Hit Database Source	5									-N		i.k	Z	IN CALL	EST HUMAN	EST HUMAN	Z	Į.	Z		Į.	FIN	E P	LN		EST HUMAN	١.	- LN	EST LIMAN	NAM IL FOR	-1	-N-	EST HUMAN	- 22
Single	op Hit Acession No.	F240786 1		18002R2	TN 28282	AI 163252 2	0718	8400716INT	TN 88888N	TA SERIES NT	A DO 4 0 2 2 4		-		1	1	AW193024.1	6912457 N I	6912457 NI	32095	5.		T	T	AF273041.1	AFZ/3041.1	1100142	720556 NT	3939077	1 / / / / / / / / / / / / / / / / / / /	AA077589.1	AA077689.1	(65/406 NI	747900	
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	Expression Signal	1	4.27	1.35	2. 3	1.04	2 3	3.08	B) (2.40	2.49	1.36	1.36	1.69	1.69	1.57	1.57	5.96	5.96	2	1.19	1.58	1.58	3.75	1.02	1.02	1.64	4.13					2.41		0.97
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	SEQ ID		14670	14675	14684	14684	14695	14697	14697			14709	14709	Ι.	L	L	┸	1_	١.	1_	14721	14722	3 14722	14729	14749	14749	3 14779	14785	3 14155	3 14155	5 14787	5 14787	7 14789		0 14792
	Probe SEQ ID NO:		1935	1940	1949	1949	1959	1961	1961	1962	1962	1973	1973	1979	1979	1981	1981	1982	1982	1984	1985	1986	1986	1993	2014	2014	2046	2052	2053	2053	2055	2055	2057	2059	3060

Page 484 of 536 Table 4 Single Exon.Probes Expressed in Brain

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olingia Exolitriodes Expressed in Brain	Top Hit Descriptor	qv30f08.x1 NCI_CGAP_Ut2 Homo sapiens cDNA clone IMAGE:1888871 3' similar to contains Alu repetitive element:	80148514651 NIH MGC 68 Home sentens cDNA clara IMA CE 20007777 51	601902604F1 NIH MGC 19 Home seniens of NA close MA of CA 1492322 E1	601902604F1 NIH MGC 19 Home sentens cDNA clara MA CE:4138222 5	RC3-CT0413-270700-022-d10 CT0413 Home saniens cDNA	RC3-CT0413-270700-022-d10 CT0413 Homo Banians cDNA	Human plasma membrane calcium ATPasa Isoform 2 (ADT202) mDNA	Human plasma membrane calcium ATPase (soform 2 (ADT2R2) meNA combis cas	Homo saplens GTP binding protein 1 (GTPRPA1) mRNA	QV1-GN0065-140800-318-c10 GN0065 Home seniens cDNA	Homo sabiens X-linked tuvenille retineschists meden (1854) gans ages 6 and annual statements	601672066F1 NIH MGC 20 Home seniers china clare MAA CE-3064755 5:	PM0-BT0547-210300-004-F04 BT0547 Homo sepiens CDNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	901.04	Homo explains mathematical control to the control of the control o	Troitio seprens interaction opinionale receptor 1 aprile (mGluK1alpha) mRNA, complete cds	QV-BT065-020399-092 BT065 Homo saplens cDNA	Homo saplens potassium large conductance calcium-activated channel, subfamily M, beta member 3-like KCNMAR3) mRNA	Human DNA-binding protein mRNA 3'snd	601122338F1 NIH MGC 20 Home sabiens cDNA clone IMAGE 3346688 E	AV738288 CB Homo saplens cDNA clone CBNRDE08 5'	AV738288 CB Homo saplens cDNA clone CRNRDEOR F	0032e01.81 NCI CGAP Lu5 Homo sanians chiva clara IMA CE11567606 91	602014829F1 NCI CGAP Bm64 Homo seniene chNA clone IMAGE:446724 E	601572186T1 NIH MGC 55 Home sapiens cDNA clare IMAGE: 22300132	CM1-TN0141-250900-439-b08 TN0141 Homo sentens chula	CM1-TN0141-250900-439-b08 TN0141 Homo saciens CDNA	601900281F1 NIH MGC 19 Homo sapiens cDNA clone IMAGE 4128822 F	bb84e02.yf NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3049082 5' similar to TR:Q15170 Q15170 TRANSCRIPTION FACTOR S-II-RELATED PROTEIN;
ולוים באטוו רוס	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN		LZ LZ	NT	EST HUMAN	N	EST HUMAN	EST HUMAN	F	FST HUMAN	ч	EST HIMAN		Į į	Į.	EST HUMAN	EST_HUMAN	EST HUMAN	EST. HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	Т	Г	
5	Top Hit Acession No.	1244247.1		0.0E+00 BF315325.1	0.0E+00 BF315325.1	0.0E+00 BE697125.1	0.0E+00 BE697125.1	.00620.1	.00620.1	4758489 NT	0.0E+00 BE767964.1	0.0E+00 AF018963.1	F027562.1	0.0E+00 BE072624.1						7657252 NT	14787.1	3.1	Г				48899.1			Γ	
	Most Similar (Top) Hit BLAST E Value	0.0E+00 AL	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 L00620.1	0.0E+00 L0C	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+001A	0.0E+00	0.0F+00 78827 4	0.0E+00 A 904640 1	0.0E+00 AI904640.1	0.0E+00	0.0E+00 L14787.1	0.0E+00 BE	0.0E+00 AV738288.1	0.0E+00 AV738288.1	0.0E+00 AA931691.1	0.0E+00 BF344434.1	0.0E+00 BE7	0.0E+00 BF377897.1	0.0E+00 BF377897.1	0.0E+00 BF313617.1	0.0E+00 BE018750.1
	Expression Signal	1.78	3.46	1.48	1.48	3.07	3.07	3.71	3.71	1.36	2.06	1.59	3.76	1.03	1.06	1.3	1.47	6.39	6.39	1.05	1.22	1.05	7.59	7.59	1.4	5.68	12.14	3.55	3.55	2.04	1.93
	ORF SEQ ID NO:		27526	27528			27533	27544		27549			27572	27573	27574	27576	27577	27579	27580			27644	27647	27648	27650	27654	27655	27659	27660	27665	27668
	Exon SEQ ID NO:	14784	14799	14801	14801	14804	14804	14812	14812	14817	14838	14839	14841	14842	14844	14847	14848	14850	14850	14883	14908	14912	14914	14914	14916	14920	14921	14925	14925	15588	14931
Į.	Probe SEQ ID NO:	2062	2067	2069	2069	2072	2072	2080	2080	2086	2107	2108	2110	2111	2113	2116	2117	2119	2118	2153	2179	2183	2185	2185	2187	2191	2192	2196	2196	2200	2203

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	Top Hit Descriptar	JASSON 81 Spares, pregnant uterus, NbHPU Homo sapiens cDNA done IMAGE: 486540 3' similar to	gb:X68857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN); gb:X68857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	2453-07.s1 Soares_pregnant_uterus_Nahr_C1 toning_cape. gb:X65857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	Homo saplens chromosome 21 segment HS21C004	Homo saplens chromosome 21 segment HS21C004	Homo sapiens KIAA0952 proteth (KIAA0952), mKNA	Homo sapiens KIAA0952 protein (KIAA0952), mRNA	Human beta-prime-adaptin (BAM22) gene, exon 16	Homo sapiens E1A binding protein p300 (EP300) mRNA	Homo sapiens KIAA0952 protein (KIAA0952), mRNA	R01433525F1 NIH MGC 72 Homo saplens cDNA clone IMAGE:3918607 b	PAYAGE ANH MGC 70 Homo sapiens cDNA clone IMAGE:3897457 5	SOLVESTARET NIH MGC 70 Home saplens cDNA clone IMAGE:3897457 5	Home sapiens mRNA for KIAA1363 protein, partial cds	Home saniens differentially expressed in FDCP (mouse homolog) 6 (DEF6), mRNA	Truitio septembrilla in the septembrilla of th	Hornic septials university and intermediate special septials and septials and septials and septial septials and septials are septials and septials and septials and septials and septials are septials and septials and septials are septials and septials and septials are septials and septials and septials are septials and septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials and septials are septials are septials.	CAUSOU XI Source total fetus Nb2HF8 9w Hamo saplens cDNA clone IMAGE:759740 5	TY 2011 1 Source: Intel fetus Nb2HF8 9w Homo sapiens cDNA clone IMAGE: 739740 5	ANSTRUCTURE INC. CGAP_Brn67 Homo saplens cDNA clone IMAGE:4157339 6	Homo sapiens potassium channel Kv2.1 mRNA, complete cds	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens flavin containing monooxygenase 3 (FMUs), mr. No. 2007 21 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	7722a02.x1 NCI_CGAP_CLT1 Homo saplens cDNA clone IMAGE:3289370 3 SImilia (2 11.00-100-100-100-100-100-100-100-100-10	KIAA0857 PROTEIN;	Homo sapiens prosping area or made or	ty67008.x1 NCI_CGAP_UZ home selfers con a care and a care and a care and a care	Homo saplens KIAA0952 protein (KIAA0952), minuk	Homo sepiens KIAA0952 protein (KIAA0952), mKNA	Homo saplens sperm specific antigen 2 (SSFA2), mKNA	Homo saplens sperm specific antigen 2 (SSFA2), mkNA	Homo sapiens KIAA0218 gene product (KIAA0218), mRNA	
	Top Hit Database Source		EST_HUMAN	FST HUMAN		TZ.	5		L L	1	1	Not all For	HOLL HOMAN	HOLL HOMEN	ESI HUMAN	Z.	ž	Ę	EST HUMAN	ESI HOMAN	EST HUMAN	FOI TOWN	5	7	Į.		EST_HUMAN	۲	EST HUMAN		LN L	TN	TIVE	TN Z	
26	Top Hit Acession No.		AA042813.1			T	7007404 MT	1002401	1007#700/	036204.1 AEE7EE NT	4937 330	1862401 IN	BE895281.1	BE905563.1	BE905563.1	AB037784.1	11545748 NT	11545748 NT	A1076404.1	AA429001.1	AA429001.1	BF347039.1	LU2840.1	AB020/17.1	ABOSO IIII	02520	0.0E+00 BE676095.1	0.0E+00 AF044571.1	A1625542.1	7882401 INT	TM 104CART				
	Most Similar (Top) Hit BLAST E	value	0.0E+00 A	l d	0.05-00	0.0E+00.P		0.0E+00	_		0.0E+00	0.0E+00	0.0E+00 E		0.0E+00		0.0E+00	0.0E+00			_		0.0E+00	0.0=+00	0.05+00	1							0.05+00	1	
	Expression Signal		£.		1.55	3.37	3.37	2.3	2.3	1.37	5.71	2.03	1.71	1.27	1.27	2.35	4.02	4.02	1.87	2.33	2.33	1.75	1.52	1.61		1.39	117								1.24
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Single Exon ORF SEQ Expression Most Strniler Top Hit Accession Deabase Source 15058 27794 1.24 0.0E+00 E774678 NT Top Hit Accession Deabase Source 15058 27794 1.24 0.0E+00 E774678 NT EST HUMAN 15077 27813 2.96 0.0E+00 E7407076.1 EST HUMAN 15077 27813 2.96 0.0E+00 AV887076.1 EST HUMAN 15078 27814 2.03 0.0E+00 AV887076.1 EST HUMAN 15081 27814 2.03 0.0E+00 AV887076.1 EST HUMAN 15081 27814 2.03 0.0E+00 AV887076.1 EST HUMAN 15082 27816 27816 2.03 0.0E+00 AV887076.1 EST HUMAN 15082 27816 2.98 0.0E+00 AV887076.1 EST HUMAN 15082 27820 0.98 0.0E+00 AV887076.1 EST HUMAN 15082 27820 0.98 0.0E+00 AV887076.1 EST HUMAN 15182 27872 1.06 0.0E+00 AV887076.1 EST HUMAN 15184 27872 1.06 0.0E+00 AV887076.1 EST HUMAN 15184 27872 1.06 0.0E+00 AV88708.1 EST HUMAN 15184 27872 0.98 0.0E+00 AV88708.1 EST HUMAN 15184 27872 2.09 0.0E+00 BE85605.1 EST HUMAN 15184 27872 2.09 0.0E+00 BE85605.1 EST HUMAN 15184 27872 2.09 0.0E+00 BE85605.1 EST HUMAN 15184 27827 1.22 0.0E+00 BE85103.1 EST HUMAN 15184 27827 1.22 0.0E+00 BE83103.1 EST HUMAN 15184 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27827 0.0E+00 BE83103.1 EST HUMAN 15188 27	bes Expressed in Brain	Top Hit Descriptor	Homo saniene KIA40219 mani mad in 1/KIA 400.00	Homo contain plant in the product (NIMAUZ18), MRNA	A 1134142 NTODB2 U	POTSERVER IN ZINES TOTAL Septens COINA CIONE IN I ZRP3002064 5	MR4.SND033.430400.042.4.701400.1.1	Homo certiene KIA \$0244 ===================================	Home contain history of the protein (KIAAU244), mKNA	Home seriens havese-o-prosporate dehydrogenase (glucose 1-dehydrogenase) (H6PD), mRNA	Home series successive DAFO, mRNA	CONTROL OF STATE CONTROL OF STATE CONTROL OF STATE CAST CONTROL OF STATE CAST CONTROL OF STATE A (CYP3A4) and cytochrome P450 polypeptide 7 (CYP3A7) genes. complete cds: and cytochrome D450	polypeptide 5 (CYP3A5) gene, partial cds	AU118082 HEMBA1 Homo saplens cDNA clone HEMBA1002839 6	AU118082 HEMBA1 Homo sablens cDNA clone HEMBA1002820 C	AU118092 HEMBA1 Homo septens CDNA clone HEMBA1002020 5	Homo saplens hypothetical profess 120084 12 120084 1	MRO-RNOTO-APPENDED TO BE ALCOHOL (FLOCUSE) THINA	AU119582 HEMBA1 Home series about alexanders con A	ox60502.x1 Scares_NH-MIDL_S1 Homo septens cDNA clone IMAGE:1660683 3' similar to TR:O08662	Homo contain throughout a training of the contai	601432808E1 NIH MGC 72 Home company PNN - WAS BOUNDED	MR1-TN0021-280800-001-H08 TN0021 Home 5-2018	AB005622 Helle CDNA (T. Nome) Home College CDNA	Homo sablens alutamate receptor ignofrance N marked by a marked by the sablens alutamate receptor ignofrance N marked by the sablens alutamate receptor ignorpance.	Homo sapiens dene for cholecysthkinin hom A recentary consists.	Homo sablens dene for cholescenting the A security of the contraction	Homo sablens imminoriohiliha ilika francashi 1 2000 1 1 1 1 2 1 2 1 1 2 2 2 2 2 2 2	502018058F1 NCI CGAP Brigg Hams conjusted to Variant + (IL.110) gene, exch 6	domo septems collegen the XII dube 4 (2014 1984 1984)	CMO-MT0033-150870. 428-641 MT0033 LASS LASS	OMO-MT0033-150600-428-h11 MT0033 U.S.	Tuman Gordein-chinaled receptor (CDDA)	Tuman G protein-coupled recentor (CPRA) gene, complete cas	802184558T1 NIH_MGC_42 Homo sapleris cDNA clone IMAGE:4300383 3'
Exon NO: ORF SEQ ID NO: Expression Signal Most Similar PLASTE Most Similar Value Top Hit Acess ID NO: 15058 27794 1.24 0.0E+00 7892 16074 27807 3.2 0.0E+00 7892 16076 27817 2.39 0.0E+00 AU131142.1 16076 27812 3.2 0.0E+00 AU131142.1 16078 27813 2.99 0.0E+00 AU131142.1 16078 27814 2.03 0.0E+00 AV18082.1 16078 27816 2.03 0.0E+00 AV18082.1 15081 27816 2.03 0.0E+00 AU18082.1 15081 27816 2.03 0.0E+00 AU18082.1 15082 27816 2.03 0.0E+00 AU18082.1 15081 27876 7.98 0.0E+00 AU18082.1 15082 27876 0.98 0.0E+00 AU1808.2 15089 27872 0.98 0.0E+00 AU1808.2 1514	gle Exon Pro	Top Hit Database Source	LZ	μZ	EST HIMAN	EST HIMAN	EST HIMAN	L		F				EST_HUMAN	EST_HUMAN	EST_HUMAN	þ	EST HUMAN	ST HUMAN	NAMILIA FO	- L	ST HUMAN	Т	Т	Т	Т			HUMAN		Г	Т			T_HUMAN
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Exam NO: ORF SEQ ID NO: Expression 15068 27794 1 16074 27811 2 16074 27814 2 16077 27818 2 16078 27819 2 16079 27819 2 16079 27819 2 16079 27819 7 16079 27819 7 16081 27819 7 15081 27819 7 15082 27819 7 15083 27872 0 15138 27873 0 15140 27896 2 15140 27897 4 15160 27897 4 15183 27827 1 15183 27927 1 15188 27928 0 15188 27928 0 15189 27929 0 15189 27928 0 <tr< td=""><td></td><td></td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td></td><td>l</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00/</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00 B</td><td>0.0E+00 B</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00 D</td><td>0.0E+00 D</td><td>0.0E+00 A</td><td>0.0E+00 B</td><td>0.0E+00</td><td>0.0E+00 BI</td><td>0.0E+00 BI</td><td>0.0E+00</td><td>0.0E+00</td><td>0.0E+00 B</td></tr<>			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		l	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00 D	0.0E+00 D	0.0E+00 A	0.0E+00 B	0.0E+00	0.0E+00 BI	0.0E+00 BI	0.0E+00	0.0E+00	0.0E+00 B
Exan SEQ ID ID ID ID ID ID ID ID ID ID ID ID ID		Expression Signal	1.24	3.2	2.39	4.31	1.51	2.99	2.03	2.03		i t	5.26	88:/	7.98	7.98	0.98	0.91	1.06	3.74	0.98	4.44	1	86.0	4.5	2.09	2.09	1.91	1.22	4.45	1.24	1.24	0.93	0.93	2.98
		ORF SEQ ID NO:		27807	L			27813	27814	27815			77047	/10/7	27818	27819	27820		27872		27873		27878		27892	27896	27897	27906	27911	27919	27822	27923	27927	27928	27929
SEC ID NO: 2334 2335 2335 2335 2335 2355 2355 2355			Ш									15070	L		⅃	- 1	J	- 1		16137	15138	15141	15145	15154	15157	15180	15160	15168	15172	16179	15183	15183	15188	15188	15189
<u> </u>		Probe SEQ ID NO:	2334	2348	2352	2353	2364	2355	2356	2356		2357	2350	8000	8000	2359	2360	2377	2415	2416	2417	2420	2424	2433	2437	2441	2441	2449	2454	2481	2466	2465	2470	2470	2471

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																								200246					13'					
	Top Hit Descriptor	hendshot x1 NCI CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2872759 3'	UI-HF-BP0p-als-c-07-0-UI.r1 NIH_MGC_51 Homo saplens cDNA clone IMAGE-3072133	Homo sapiens protein kinase, AMP-activated, alpha 2 catalytic subunit (PRKAA2) mRNA	Homo saplens protein kinase, AMP-activated, alpha a catalyte successive succe	RC3-ST0197-300300-016-c04 ST0197 Homo Septems CONA	601592530F1 NIH_MGC_7 Homo sapiens cDNA cione IMAGE. 35700 13 C	Homo saplens death receptor 6 (DR6), mixing	UI-H-BI4-aoz-b-08-0-UI.s1 NCI_CGAP_Subs Home (XK gene)	Homo sapiens mRNA for memorane danisport processor (PDGFRL) mRNA	Homo sapiens platelet-denived grown racio, rong and IMAGE:3905148 5	601503356F1 NIH MGC_70 Home septens obtained pow121 (POM121L1), mRNA	Homo sapiens similar to rat integral membrane grycopaction of the many	Homo sapiens hypothetical protein FLU20366 (FLU20369), mixton	Human Sec62 (Sec62) mRNA, complete cus	601508211F1 NIH_MGC_71 Homo saplens cDNA clone (MAGE: 3891371 6)	601489241F1 NIH_MGC_69 Homo sapiens CDIVA clure IMAGE:3891371 5	601489241F1 NIH MGC 69 Homo sapiens dulya civile living contraction	Homo sapiens adlican mRNA, complete cds	601064738F1 NIH MGC 10 Hama Saprens CONTROL 1	Т	AU143277 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo sapiens curia cidile 11 april 2017 Y79AA1 Homo cidile 11 april 2017 Y79AA1				\neg	Т	1	Homo sapiens mRNA for KIAA1415 protein, parusa con proteins cDNA clone IMAGE:3070631 3	1	$\overline{}$	Г	$\overline{}$	
	Top Hit Database Source	- 1	EST HOMAN		FZ	EST HI IMAN	EST HUMAN	LN	EST HUMAN	LN	LN	EST HUMAN	LN	LN L	LN	EST HUMAN	EST HUMAN	EST HUMAN	LN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN		EST_HUMAN	EST HIMAN	LZ	LV	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	
Sup.	Top Hit Acession · No.		Ī	AW501010.1	TARROBEINT	45555	AW813833.1	TARZOZA NI	RFE09482.1	732684.2	5453871 NT	DE ON 037	TAK746R NT		00000	U93239.1	BE880480.1	BE0/0011.1	AES/5011.1	DEE28024 1	A11443277 1	0.0E+00 A01+0277.1	0.0L-100 AC 1	0.0E 100 BF 292896.1		0.0E+00 BF223041.1	0.0E+00 AF245505.1	0.0E+00 BEZ90613.1	0.0E+00 AB037838 1	DEE43835 1	0.0E+00 Br313630.1	DE204134 1	0.0E+00 Br 204131.1	
	道士川	_		0.00+00	0.0=+00	0.0E+00		0.05+00	0.01+00	00-10-0		0.00	0.05+00	0.01.400	0.0=+00	0.05 +00	1		0.0E+00	0.05	0.01	1		1				-		1		U.UET	0.00	
	Expression Signal		3.34	2.99	0.91	0.91	1.66	5.22	4.6	1.48	23.	3.28	1.3	1.96	2.01	2.21	1.44						7		0.1								1.83	
	ORF SEQ 10 NO:		27940	27941	27958	27959		27971	27383	27972	27974		27977		27979									١	3 28015	7 28016				11 28061			28069	١
	0	ö	15200	1.	15215	L		15231	1			15236	L	1	15240	١.	<u> </u>	L	١.	15251	<u> </u> _				16276	15277	L.	L	06 15531	15531	1_	111 15323	1 1	2615 15326
	Probe SEQ ID	Ö	2482	2484	2498	2498	2509	2514	2515	2516	2518	2520	2522	2523	2524	2525	2530	2534	2634	2536	2554	2561	2561	2562	⁵⁸	2663	2566	2589	2606	2606	2607	2811	8	²⁸

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Table 4
Single Exon Probes Expressed in Brain

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Top Hit Descriptor	br19b08.x1 NCI_CGAP_Brn25 Homo sapiens cDNA clone IMAGE:2168055 3' stmllar to gb:L20977 CALCIUM-TRANSPORTING ATPASE PLASMA MEMBRANE, BRAIN ISOFORM 2 (HUMAN): TATA to be binding norbing (TRP) associated factor, RNA polymerase II, I, 28kD (TAF2I)	MRNA	HOMO Septems mining for MAR 1439 process, parties and CE:3944304 5	60139010671 NIT MICC 7 Home septems CDNA clone IMAGE:3844304 5	601 090 1091 INIT MICC 15 Home saciens cDNA clone IMAGE:3051389 5	60 145/222 FINIH MIGG 7 Homo sepiens CDNA clone IMAGE:3939222 6	Usus series: IMP (Inosine monophasphate) dehydrogenase 1 (IMPDH1) mRNA	Holling saperis in (income holling the second by the second protein the second protein the second protein the second protein the second protein the second protein the second protein the second protein the second protein the second protein the second the	Hamp sapells bruch a shown complete data (L44L) and FTP3 (FTP3) genes, complete data	Homo septens guanyfate oyotase activating protein 4 (5000 to) garto, control of the control of t	AU133386 N12KP4 Homo sapiens duna cigie in 12K1 400 004 0	Human bullous pemphigoid antigen (BFAG1) minna, compass cus	AU130403 N IZRES Hamp septems control clare NT2RP3000779 5	ACTIONAL NI STATE OF THE SECTION OF	RC1-O10086-Z20300-011-d07 O10000 Holling septems 30000 3/	ACCOUNTY INC. CON LONG Seniors CDNA clone IMAGE:3628923 6	601236714F1 NIT MICC 30 Home series CDNA close IMAGE:3610267 5	EST 188414 HCC cell line (matastasis to liver in mouse) II Homo sapiens cDNA 6' end similar to ribosomal	protein L29	1601589625FT NIM _MCC / Tollio sapietis control money for the co	Human betta-printe-tatapun (BANVZZ) genesian wanan SMDF mRNA	Homo septens neureguint 1 (MNO 1), various production (MNO 2) pene, complete cds	Truin september of the company of th	Home saprens indexed in the Control of Services (NAGE:3945983 5)	BANGSER NIM MGC 39 Home septems CDNA clone IMAGE:3689564 5'	AVIZAGATUTE Home caniens cDNA clone HTBBYE09 5'	HV72104/ FILD Truit agricultures associated PD1 (KIAA0767) mRNA	Home sapiens spermatogenesis associated PD1 (KIAA0757) mRNA	Homo sapiens twonheited protein FLJ20477 (FLJ20477), mRNA	
Top Hit Database Source	EST_HUMAN		Т	Т	7	Т	HOMAN	Ž	. TN	NT	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	ESI HOMAN	EST_HUMAN	EST HUMAN	NT	LN L	Z	IN	ESI HOMAN	EST HUMAN	EST HUMAN	1110	L L	1212
Top Hit Acession No.	AI571737.1	5032150 NT	AB037859.1	BE795445.1	BE795445.1	BE293328.1	BE792472.1	4504686 N1	U78027.1	AF173227.1	AU133385.1	M69225.1	AU130403.1	AU130403.1	AW887015.1	BF000018.1	BE383165.1	BE531263.1	0.0E+00 AA316723.1	0.0E+00 BE794884.1	0.0E+00 U36253.1	0.0E+00 7669517 NT	AF110763.1	0.0E+00 AB051828.1	0.0E+00 BE796376.1	0.0E+00 BE563433.1	AV72164			8923441 N
Most Similar (Top) Hit BLAST E Value	0.0E+00/	_					0.0E+00	0.0E+00	0.0E+00	0.0E+00	00+30'0		i	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00				- 1	0.0E+00	╛
Expression Signal	2.08	2.19	4.95	1.02	1.02	2.55	4.92	1.51	1.27	6.55	1.22	1.08	2.22	2.22		1.25		2.57	4.21	5.57	3.83		1.78	1.27		3.48	1.18			1.21
ORF SEQ ID NO:	28072	28073	28077	28078	28079	28080		28096		28103			28109	28110			Ì_			28169	28177		28179	28181	28187	28191			28195	
Exon SEQ ID NO:	15329	15330	1_	1	15334	Ľ		15352	15381	1	1	1	L	L	Ļ	15376		15378	15432	1_	1_	L	15441	1	l	15604	L	15453	15453	15454
Probe SEQ ID NO:	2618	2819	2621	2622	2622	2625	2632	2841	2854	2852	2658	2659	2661	2681	2863	2666	2867	2668	2726	2728	2732	2733	2734	2736	2742	2745	2746	2748	2748	274

PCT/US01/00667

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Page 489 of 536 Table 4 Single Exon Probes Expressed in Brain

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Top Hit Descriptor	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo saptens hypertension-related calcium-regulated gene mRNA, complete cds	AV651068 GLC Homo sapiens cDNA clone GLCCLD07 3'	CM1-TN0141-250900-439-b08 TN0141 Homo saplens cDNA	CM1-TN0141-250900-439-b08 TN0141 Homo sapiens cDNA	Homo sapiens cerebellar degeneration-related protein (34kD) (CDR1) mRNA	Homo sapiens cerebellar degeneration-related protein (34kD) (CDR1) mRNA	601580903F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3929472 5'	Homo sapiens chromosome 21 segment HS21C001	UI-H-BW1-amw-e-07-0-UI.s1 NCI_CGAP_Sub7 Homo saplens cDNA clone IMAGE:3071340 3'	Homo sapiens chondroltin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA	Homo sapiens angiopoletin-3 (ANG-3), mRNA	Homo sapiens angiopoletin-3 (ANG-3), mRNA	602085579F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4249915 5'	Homo sapiens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA	AV725534 HTC Homo sapiens cDNA clone HTCCCA03 51	AV725634 HTC Homo sapiens cDNA clone HTCCCA03 5'	au56d04.y1 Schneider fetal brain 00004 Homo sepiens cDNA clone IMAGE:2518683 5' similar to SW:R13A_HUMAN P40429 60S RIBOSOMAL PROTEIN L13A;	602071957F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4214679 5'	601450912F1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3854642 5'	AU131494 NT2RP3 Hamo sapiens cDNA done NT2RP3002672 5'	AU131494 NT2RP3 Hamo sapiens cDNA done NT2RP3002672 5'	600944794F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:2960806 5'	600944794F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:2960808 5'	glycoprotein D=Duffy group antigen [human, blood, Genomic DNA, 3068 nt]	Homo sapiens BTRCP2 mRNA for F-box and WD-repeats protein isoform C, complete cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cds	Homo sapiens cytochrome P450, subfamily I (dioxin-Inducible), polypeptide 1 (glaucoma 3, primary Infantile) (CYP181) mRNA	Homo sapiens cytochrome P450, subfamily I (dioxin-inducible), polypeptide 1 (glaucoma 3, primary infantile) (CYP1B1) mRNA	H.sapiens serine hydroxymethyltransferase pseudogene
Top Hit Datebase Source	F	TN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	IN	EST_HUMAN	N	EST_HUMAN	LN	N	LZ LZ	EST_HUMAN	NT L	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	TN	님	IN	Ę	Į Į	NT.
Top Hit Acession No.	8923441 N	0.0E+00 AF290195.1	NO AV651066.1	0.0E+00 BF377897.1	0.0E+00 BF377897.1	4757963 NT	0.0E+00 4757963 NT	3E747193.1	0.0E+00 AL163201.2	0.0E+00 BF514110.1	4503098	7705275 NT	7705275 NT	0.0E+00 BF677694.1	7427522 NT	4V725634.1	0.0E+00 AV725534.1	0.0E+00 AI879163.1	0.0E+00 BF530661.1	0.0E+00 BE872768.1	0.0E+00 AU131494.1	0.0E+00 AU131494.1	3E300344.1	0.0E+00 BE300344.1	0.0E+00 S76830.1	AB033281.1	0.0E+00 AF264750.1	0.0E+00 AF264750.1	4503202 NT		
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00				0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00.	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00/	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 X85980.1
Expression Signal	1.21	2.5	13.89	3.13	3.13	33.8	33.8	2.58	1.15	3.36	0.99	. 2.1	2.1	4.67	1.75	13.56	13.56	7.61	5.41	5.65	1.6	1.6	20.25	20.25	4.22	4.35	8.39	8.39	3.52	3.52	3.52
ORF SEQ ID NO:	28197	28198		28199	28200	28203	28204	28209		28219		28227	28228	28229	28237	28239	28240		28245	28246	28247	28248	28249	28250			26144	26145		ĺ	28258
Exon SEQ ID NO:	15454	15455	15456		15457		15461		15476	15477		15489	15489		ŀ	15499		15501	ı	15505	ı				!		13491	13491	13776		15608
Probe SEQ (D NO:	2749	2750	2751	2752	2752	2756	2756	2760	2771	2772	27.78	2784	2784	2785	2791	2794	2794	2796	2799	2800	2802	2802	2803	2803	2809	2812	2818	2818	2822	2822	2840

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	Top Hit Descriptor	Homo sepiens 5-eminoley ilinete synthese 2 (AI AC2)	Homo saniens mRNA for KIA A 1577 States	Homo saniens chromosome 21 segment DS2 Con	Human sodium channel mRNA	Human AHNAK nucleoprotein mRNA 5. and	PM0-H70343-281299-003-602 H70343 Home septem - DNA	PMO-HT0343-281299-003-602 HT0343 Home september ANIA	H. sapiens Id3 gene for HI H trae transmission forthan	Homo saplens chromosome 21 segment MS21Chas	Homo saplens zinc finger profess 734 (ZNE234)	Homo saplens zinc finger protein 224 (ZNE224) mDNA	Homo sabians zinc finaer protein 224 (ZNE224) mDNA	Homo sapiens camma-cytoplasmic actin (ACTODs) security.	Homo sapiens gamming-ovinglesmic actin (ACTGP3) pseudogene	Novel human mRNA from chromosome 1 which has aligned by the party of t	Hisapiens mRNA for nuclear DNA halicase II	Homo saplens protocadherin alpha C1 (PCDH-alpha C1) mBNA complete c1.	Homo saplens eukarvolic translation elongation factor 4 alaka 4 (EEE444) 2014	Homo seplens eukaryotic franslation elongation factor 1 ainha 1 (EEF 41) mbNA	Homo saplens serine/threonine kinase 9 (STK9) mRNA	DKFZp688G0621 r1 686 (synonym: hute1) Home saniens cDNA clare DK E2288820824	Homo sapiens KIAA0054 gene product Helicase (KIAA0054) mRNA	Homo sapiens KIAA0054 gene product: Helicase (KIAA0054), mRNA	Homo saplens chondroitin sulfate proteoglycan 4 (melanoma-associated) (CSPGA) mRNA	QV2-BT0636-130400-138-h03 BT0636 Homo sapiens cDNA	QV2-BT0636-130400-138-h03 BT0636 Homo sabiens cDNA	Homo sapiens chromosome 21 seament HS/1Cons	Homo saplens chromosome 21 segment HS/1C006	Homo sapiens hHb5 gene for hair keratin expns 1 to 9	Homo saplens EphA4 (EPHA4) mRNA	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEE1441) mBNA	b18d07.x1 NCI_CGAP_Bm25 Homo saplens cDNA clone IMAGE:2167981 3' similar to TR:O16247 O16247 F44E7.2 PROTEIN ;
2.6	Top Hit Database Source	LZ.	I-N	L Z	Įγ	Ł	EST HUMAN	EST HUMAN	Ļ	Ā	Ę	Ĭ	Ę	Ľ	N-	N-	Z	LN L	Ę	Ę	Ę	EST_HUMAN	LN	N _T	TN	EST_HUMAN	EST_HUMAN	N	N	Į.	F	F	EST_HUMAN
	Top Hit Acession No.	+00 AF068624.1	+00 AB040960.1	AL 163201.2	0.0E+00 M91803.1	+00 M80902.1	0.0E+00 BE154504.1	+00 BE154504.1	0.0E+00 X73428.1	AL163268.2	0.0E+00 7019584 NT			D50657.1	D50657.1	0.0E+00 AL096857.1	Y10658.1	+00 AF152303.1	4503470 NT	4503470 NT	4507280 NT	+00 AL047599.1	7661883	7661883 NT	33098				H00 AL163206,2		4758279 NT	4503470 NT	00 AI561002.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D50657.1	0.0E+00	0.0E+00	0.0E+00 Y10658.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 ₽	0.0E+00 \	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.34	1.22	2.61	263	1.94	1.29	1.29	1.18	2.59	1.03	1.03	1.03	18.48	18.48	1.69	7.2	1.42	47.87	47.87	2.35	1.06	0.97	0.97	2.96	5.46	5.46	2.09	2.09	4.18	1.33	20.94	1.91
	ORF SEQ ID NO:			28263							28272		28274	28277	28278	28281			28282	28283	28292	28296	28297	28298		28300	28301	28312	28313		28321	28322	28323
	σ	15609	15611	15617	15620	15622					15629	- 1			1	15636		- 1	- 1	15639	15849	15652	15853	15663	15654	15657	15657	15864	15664	15671	15673	15674	15675
	Probe SEQ ID NO:	2841	2843	2849	2862	2854	2857	2867	2859	2860	2861	2861	2861	2866	2866	2869	2870	2871	2872	2872	2882	2883	2886	2888	2887	7880	2880	2897	2897	38 38 38	2807	88	2909

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	Top Hit Descriptor	Homo sapiens semenogalin I (SEMG1) mRNA	Homo sapiens membrans-bound aminopeptidase P (XNPEP2) gene, complete cds.	Home conjens host chrok 70kD protein 1 (HSPA1A), mRNA	mo opposite the state of the st	Homo septems near snock / u/u/ protein (rior Ara), makes	Isoform 2 of a novel human micha from chromosome 22	Home saplens putative transcription factor CR63 (CR63) mRNA, partial cus	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, 164 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaprophysin genes.	complete cds; and L-type calcium channel a>	Homo sapiens chioride channel CLC4 (CiC4) mrNN, compare cus	Human germine gene 16.1 for ig lannous L-cliain O legion (195 - 01.0.)	Homo sapiens F-box protein FBL5 (FBL5) mRNA, complete cas	Homo sapiens melanoma-associated antigen (MAGE-C1) gene, complete cus	Homo sapiens SWI-SNF complex protein p2/0 mKNA, partial cas	Homo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo saplens KIAA0469 gene product (KIAA0409), mkiva	Homo saptens olfactory receptor-like protein (OLFR 42B) gene, OLFR 42D-9110 away, pai uai ous	Homo sapiens potassium voltage-gated channal, Shab-related subfamily, member 1 (KCNB1) mRNA	Human feritin heavy chain mRNA, complete cds	Homo sepiens mRNA for KIAA0549 protein, partiel cds	Homo saplens mRNA for KIAAU349 protein, partei cus	yesztős at Stratagerie lung (res/ 2.10) normo sapram corry como marchine. S28639 BASIC PROTEIN, 23K - ;	601878507F1 NIH_MGC_55 Homo sapiens cDNA clone IMACE: 4107433 5	H.saplens mRNA for gamma-guriamytransferese	H.sapiens mkina for garming-grutarity a reference	Homo sapiens neurexin III (NRXNs) mRNA	Homo saplens neuraxin III (NKXN3) mxNA	Homo saplens interleukin 1 receptor, type I (ILTR1) mRNA complete cds	Homo sapiens nucleolar phosphoprorein 623 (NPM / Innvv.), compress cas	Home sapiens solute carrier raminy 23 (minochondrial protein, mana). (SLC25A5), nuclear gene encoding mitochondrial protein, mRNA	Homo sapiens CREB binding protein (Rubinstein-Taybi syndrome) (CREBBP) mrwA
-	Top Hit Database Source							NT	¥ 5				Ĭ	NT	NT IN							Į.		EST_HUMAN 0	-					Į,		
	Top Hit Acession No.	4506882 NT		0.0E+00 AF 190955.1	1N 6946269	0.0E+00 5579469 NT								0.0E+00 AF084589.1			7662139 NT	0.0E+00 AF042075.1	4826783 NT	00 1.20941.1	00 AB011121.1	00 AB011121.1		3.1				4758827 NT	4504658	M28699.1	4502098 NT	TABOREINT
-	Most Similar (Top) Hit ELAST E	00+300	100.100	0.0=+00.0	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00 ₽	0.0E+00 X03529.1	0.0E+00	0.0E+00	0.0E+00.7	0.0E+00	0.0E+00	0.0E+00	0 05+00	O.OE+	0.0E+	0.0E+	0.0E+00 T94870.1	0.0E+00	0.0E+	0.0E+00 X98922.1	1	1			0.0E+00	
	Expression Signal	27.0		4.62	3.74	3.74	5.88	18		1.98		2.8		1.72	3.2						1.32										1.98	
	ORF SEQ ID NO:	200446	01407		28454	28455		28480	1			28483		28490			28514		_			28560		28584			L	28802				
	Exon SEQ ID NO:	- [16808	į.	ı	l	1	15818	┸	L	1	L	L	.L.	15875	1		1591	1.	L	<u> </u>	1	1	ı	L	┸	L	1	1	1
	Probe SEQ ID NO:		303	3038	3042	3042	3044	3	9400	3052	3066	3074	1000	2000	200	3105	3 5	2444		3138	3151	3161	0460	3172	3178	3478	3187	2467	346	3211	1 8	3214

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Top Hit Descriptor	Homo sapiens CREB binding protein (Rubinstein-Laya) syndronia) (CNEDD) / 112.00.	ae87b11.s1 Stratagene schizo prairi 311 monto appeara	Homo sapiens anglostatin binding protein 1 mRNA, complete cas	Homo sapiens angiostatin binding protein 1 mRNA, complete cds	Homo sapiens fibrillin 1 (Marfan syndrome) (FBN1) mRNA	Homo sapiens titln (TTN) mRNA.	Human connectin 43 processed pseudogene	Homo saciens HLA class III region containing tenascin X (tenascin-X) gene, partial cds, cytochrome France.	hydroxylase (CYP21B), complement component C4 (C4B) G11, helicase (SKIZW), KD, complement of hydroxylase (SYIZW), KD, complement component C2 (C3) denes.>	(bt), and complement compared to the complete ods. Howevery large G-protein coupled receptor-1 (VLGR1) mRNA, complete ods	Home saniens Kladoddo protein (Kladoddo), mRNA	Home seniens KIA 40440 protein (KIAA0440), mRNA	Houng seniens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	House series A triagge (PBKA) anchor protein 1 (AKAP1), mRNA	Indiano sapiens SWI-SNF complex protein p270 mRNA, partial cds	Turing september of the	House seniors death receptor 6 (DR6), mRNA	Aut See 18 ST NCI CGAP Brz Homo sapiens cDNA done IMAGE:1632038 3'	Home caplans G brotain-coupled receptor 24 (GPR24), mRNA	HISBROB. ZZ NCJ CGAP Pan1 Homo sapiens cDNA clone IMAGE:2222536 3' similar to SW:KL11_NAI	P25121 60S RIBOSOMAL PROTEIN L11. ;contains Alu repetitive element.	Home sapiens teromerase reverse transcriptese (TERT) gene, exons 1-6	Homo sapiers monally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sanians homonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo saniens caspase 8, apoptosis-related cysteine protease (CASP8) mRNA	Home senions cashase 8 apoptosis-related cysteine protease (CASP8) mRNA	Homo sapiens pyrin (MEFV) gene, complete cds	Homo sapiens mRNA for KIAA1507 protein, partial cds	ROTAZA995F1 NIH MGC 67 Homo sapiens cDNA clone IMAGE:3868246 5'	LANGE AND CAP GC6 Homo sapiens cDNA clone IMAGE:2305279 3' similar to TR:091929 UNIVERSITY	ZINC FILEST PROTEIN: STATE CONA clone NT2RM2000735 5'	AU123004 N LATURE TO THE SEPTEMBER OF TH	
Top Hit Database Source		T HUMAN		FIA	FIX	Į.	1			Z	Z	L	ĮN.	ĮŅ.	Į.	Z	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	NAMA LI	-1	Z	EST HUMAN	卜	Ł	IN	INI	LN .	IN 2	Į.	2	ESI HOMAN	EST_HUMAN	EST HUMAN	l I
Top Hit Acession No.	4758055 NT	A A 77 A 783 4	T	1	AF286598.1	0001004	20//20	M65189.1			AF055084.1	7662125 NT	7662125 NT	4502014 NT	4502014 NT	AF265208.1	8923624 NT	7657038 N	AA994842.1	4885312 N I			0.0E+00 AF128893.1			4502582 NT	4502582 NT	0.0E+00 AF111163.1	0.0E+00 AB040940.1	0.0E+00 BE779039.1	0.0E+00 AI632569.1	0.0E+00 AU123664.1	
Most Similar (Top) Hit BLAST E Value	00+400		_		0.0E+00 A	0.0E+00	0.0E+00	0.0E+00 M		0.0E+00 A	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.00	0.0E+00	0.0E+00	0.0E+00		0.0E+00		0.0E+00					1
Expression Signal	ממ	3	3.07	5.43	5.43	1.36	F	5.8		1.26	3.96	1.11	1.11	2.09	2.09	2.25	1.17	1.22	1.09	1.18	8				1.22			9.77	1.29	0.84	0.72		
ORF SEQ ID NO:	- 1	/2007	28638	28646	28647	28653				28668	l		L		L		L		28710		00430											١	
Exon SEQ 1D NO:		15983	15985	15993	15993	16004	16010	16017		16018	16020	16023	16023	17875	17875	Į.	1		16062	1	l	16078	1	1_	L	丄	L	L	1_	L	1	1	16130
Probe SEQ ID		3220	3222	3230	3230	3242	3248	3255		3258	3258	3284	3284	1000	3280	2000	328	3297	2300	3300		3318	2220	3327	3327	1700	338	2223	3335	2342	3	3352	SSS

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Single Exon Drobes Expressed in Busin	Top Hit Descriptor	Omo saniane offertant researcher formille 40 16 11. O	Homo capiene offsoton; receptor, remains 10, subramily C, member 1 (OK10C1), mRNA	Homo sepiens heuroblestome emplified matein // OCSESSY Homose 1 (OR10C1), mRNA	THE STATE OF THE S	Fromo sapiens 1-type cardum channel alpha1 subunit Alpha11-a Isoform (CACNA11) mRNA, complete cds	Homo septems KTA 40052 septem VKTA AAASS	Homo septions KIAAAGES protein (KIAAAGES), mKNA	Homo sapiens beaded filament structural protein 1 filancia, IBESD41 DNA	Homo sepiens leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 2 (LILRA2), mRNA	Homo sapiens skeletal muscle (Manadain 4 (EUI 4) anno annulata 4	Homo saplens death recentor 6 (DRs) mRNA	Bacteriophage P1 replication region including repA, parA, and parB genes and incA, incB, and incC incompatibility determinants	Homo sapiens profesing processes when home seement and a september of the second secon	WP14410X1 NOI_CGAP_Luid Homo septens cDNA clone IMAGE:2484819 3' similar to TR:073634 073634 NEURAL CELI A APHESION MOI ECLI II E	wp14d10.x1 NCI_CGAP_LU19 Home septens dDNA clone IMAGE:2464819 3' similar to TR:073834 073834	IONO Saplens mRNA for metallico calculario	Homo saplens W/os FBJ murha osteosarcoma viral concentral branch (FC)	Homo sapiens y-fos FBJ murine ostensarcoma vital momenta homolog (FOS), mining	Human endogenous retrovirus HERV-K10	Human MDS14 (AML1/MDS1 fusion) mRNA partial cite	Homo sapiens hypothetical protein (AE038180) mRNA	Homo sapiens hypothetical protein (AE038469), minys	Homo sapiens cell-line KG1 transcriptional regulatory protein p54 mRN4 complete c4c	Homo sapiens cell-line KG1 transcriptional requilabry protein n54 mRNA complete cuts	Homo sapiens chromosome 21 unknown mRNA	601143853F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3051373 5'	801143853F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:3051373 5'
l ford nove	Top Hit Database Source	Į.	FZ	17		EST HIMAN	-1	5	느				Ŀ		EST HUMAN	T											П	HOMAN
Sign	Top Hit Acession	7363436 NT			A E 211 100 1	V867015 1	7862401	7662401 NT	4502398 NT	F803067 NT	1076	7657038 NT	380.1	7427522	0.0E+00 Al935159.1 E			52332	6552332 NT	114123.1 NT		9558718 NT	9558718 NT	F045452.1 NT	F045452.1 NT			A0ZBC
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	00+	0.0E+00 AV	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AF1	0.0E+00	0.0E+00 K02	0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M14123.1	0.0E+00 U43293.1	0.0E+00	0.0E+00	0.0E+00 AF045452.1	0.0E+00 AF045452.1	0.0E+00 AF231922.1	0.0E+00 BE304791.1	0.0E+00 BE304791.1
	Expression Signal	0,95	0.95	1.38	1 42	0.99	1.66	1.66	1.34	5.37	1.75	2.4	5.47	1.6	3.54	3.54	2.48	1.82	1.82	1.31	5.4	1.18	1.18	1.84	1.84	1.12	2.21	0.92
	ORF SEQ ID NO:	28807	28808	28810	28811		28824	28825	28826	28827	28179	28840	28844	28846	28853	28854	28859	28868	28869	28874	28880	28884	28885	28880	28891	28898	28905	28909
	Exan SEQ (D NO:	16156		16159	16160		16175	16175	16176	16178	16441	16191	16194	16196	16204	16204	16208	16215	16215	16220	16226	16231	16231	16235	16235	16241	16252	16255
	Probe SEQ ID NO:	3398	3398	3401	3402	3406	3418	3418	3419	3421	3430	3435	3438	3440	3448	3448	3452	3469	3459	3464	3470	3475	3475	3479	3479	3484	3490	3499

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Top Hit Descriptor	te35g12.x1 Soeres_Nh-HMPu_S1 Homo sepiens cDNA clone IMAGE::2088742 3' similar to TR:000498 000498 MYASTHENIA GRAVIS AUTOANTIGEN GRAVIN ;	Human endogenous retroviral DNA (4-1), complete retroviral segment	AV701869 ADB Homo sepiens cDNA clone ADBDAH06 5'	Homo sapiens semenogelln II (SEMG2) mRNA	Homo sapiens homologous yeast 44.2 protein mRNA, complete cds	Novel human gene mapping to chomosome X	Homo sapiens mRNA for KIAA1476 protein, partial cds	Homo saplens sal (Drosophila)-like 1 (SALL1), mRNA	Homo sapiens sal (Drosophila)-like 1 (SALL1), mRNA	ox77c11x1 Sogres_NhHMPu_S1 Homo septens cDNA clone IMAGE:1682356 3' similar to WP:T19B4.4 CE13742 :	Homo seplens butyrophilin, subfamily 3, member A3 (BTN3A3), mRNA	QV0-CT0225-230300-169-e01 CT0225 Homo sepiens cDNA	Homo sepiens gamma-glutamyloysteine synthetase (GLCLC) gene, partial cds	602084583F1 NIH_MGC_83 Homo sapleris cDNA clone IMAGE:4248596 5'	QV0-DT0047-170200-123-901 DT0047 Homo sapiens cDNA	602152488F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4293643 5'	602152486F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4293645 5'	Homo sapiens retinoblastoma-binding protein 2 (RBBP2) mRNA	hi84g01.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2979024 3'	hi84g01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2979024 3'	Homo sapiens heparan sulfate (glucosamine) 3-O-sulfotransferase 1 (HS3ST1) mRNA	Homo saplens KIAA0806 gene product (KIAA0806), mRNA	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA	Homo sapiens midline 1 (Opitz/BBB syndrome) (MiD1) mRNA	Homo sapiens mRNA for G protein-coupled inward rectifier potassium channel, complete cds	Homo saplens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA	Homo sepiens WAVE2 mRNA for WASP-family protein, complete cds	Homo sapiens SH2-containing protein Nsp2 mRNA, complete cds	Homo saplens SH2-containing protein Nsp2 mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C004	Homo sapiens chromosome 21 segment HS21C004	MR2-CT0222-281099-005-e05 CT0222 Homo saplens cDNA
Top Hit Database Source	EST_HUMAN	FZ	EST_HUMAN	NT	TN	NT	LN	N	LN	EST HUMAN	Z	EST HUMAN	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	. TN	EST_HUMAN	EST_HUMAN	NT	NT	NT	TN	NT	NT	MT	NT	NT	NT	NT	EST HUMAN
Top Hit Acession No.	AI384007.1	M10976.1	AV701869.1	4506884 NT	AF078868.1	AL133204.1	AB040909.1	6997248 NT	6997248 NT	A1081907.1	325463	AW852217.1	AF118846.1	BF676393.1	AW937977.1	BF672054.1	BF672054.1	4826967	AW664693.1	AW664693.1	4826763 NT	7662319 NT	4557752 NT	4557752 NT	D87327.1	7669491 NT	AB026542.1	AF124250.1	AF124250.1	AL163204.2	AL163204.2	AW851714.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	7 =	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00			0.0E+00	0.0E+00		0.0E+00
Expression Signal	0.89	1.11	1.29	0.85	1.74	1.49	1.21	1.37	1.37	69 0	45.	4.17	0.78	6.46	6:0	0.74	0.74	0.95	1.08	1.08	1.42	0.93	0.82	0.82	2.67	33.2	2.6	3.38	3.38	1.85	1.85	1.62
ORF SEQ ID NO:	28914	28917	28940	28941		28945	28948	28963			28967	L		28976	28988	28996	28997		28999	29000	29004	29007	29015	29016	29030		29049	29051	29052	29061	29062	29065
Exon SEQ ID NO:	16260	16263	16285	16286	16287	16295	16297	16316	16316	16317	16319	L	16331	16332	16343	16356	16356	16357	16359		16362	16364	16374	16374	16391	16395	16411	16413	16413	16421	16421	16424
Probe SEQ ID NO:	3504	3507	3529	3530	3531	3539	3542	3561	3561	3562	3564	3569	3576	3577	3589	3603	3603	3604	3608	3606	3609	3611	3621	3621	3638	3642	3658	3660	3660	3668	3668	3671

Page 496 of 536 Table 4 Single Exon Probes Expressed in Brain

	Т	7		Γ	T	Τ	Τ	Τ	Ī	Ŧ	T	T	T	Τ	T	T	T	T	T	T	T	Ĭ	T	Γ	Γ	Τ	T	T	٦			Ī		Ī
Top Hit Descriptor	Homo sapiens matrix metalloproteinase 24 (membrane-inserted) (MMP24), mRNA	Homo saplens mRNA for KIAA0796 protein, partial cds	Home seriens mRNA for KIAA0910 protein, peritel cds	TIONIO SEPURIO IN UNIVERSITY OF TAXABLE SEPURIOR OF TAXABLE SEPURI	HOMO Saplens mining Idinated to protein, parta our	Homo sabiens NYANOVO proteinas (NYANOVO) ININANA	Homo sapiens KIAA0670 protein/acinus (KIAA0070), IIINNA Pomo sapiens KIAA0670 protein/acinus (KIAA0070), IIINNA	UI-H-BW0-gis-e-12-0-UI:ST NCI CCAP Substitution explains contact invocations of the contact invocation of the contact of the contact invocation of the contact of the conta	UI-H-BWU-gle-e-12-U-UI-SI INCI CONT CONDO PARTIES CONTO CONT	aa06g01rf Soares, NhhMPru, S1 Homo septens curve, cone invocation arrows comments. SW:KR84_SHEEP P02445 KERATIN, HIGH-SULFUR MATRIX PROTEIN, IIIB4. [1];	Homo sapiens mRNA for KIAA0903 protein, parual cus	Homo septiens similar to fat integral mentiorate group to the contract to the contract of the	Homo saplens mRNA for KIAA1414 protein, partial cos	Homo sepiens KIAA0569 gene product (KIAA0569), mKNA	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo saplens v-ets avian erythroblasbosis virus E.26 oncogene related (E.N.), minny	Homo sapiens v-ets avian enythroblastosis virus E28 oncogene related (EKG), mKNA	Homo sapiens DNA mismatch repair protein (MLH3) gene, complete cus	Pan troglodytes olfactory receptor (PTR208) gene, partial cds	Homo saplens similar to rat Integral membrane glycoprotein POM121 (POM121L1), mkinA	Homo sapiens similar to rat integral membrane diycoprotein POM121 (POM121CLI), minner	Homo sapiens RAB9, member RAS oncogene taminy (RAB9) minina	Mus musculus junctophilin 1 (Jp1-pending), mrsvA	(e62110x1 Soares NPL GBC 31 Turino septiatis conta complete ords	Homo sapiens protoceanerin bera 3 (r CDT-beras) III (r.c.), Compose and Compos	Homo sapiens desmopiakin (UP), UP11) (UOF) IIINNA	Homo sepiens ATP-sensitive Inwardly rectifying K-channel subunit (KCNJ9/BIR1) gene, complete cds	Homo saplens methyl CpG binding protein 2 (MECP2), mRNA	Homo saplens KIAA0589 gene product (KIAA0569), mRNA	Home seriens myosin light chain kingse Isoform 2 (MLCK) mRNA, complete cds	Library soutions mayor light chain kinase isoform 2 (MLCK) mRNA, complete cds	Thomas de Track and DWP2 complete and partial cds	Home services general or TMEMs and PWP2 complete and partial cds	מקינות מקינות מינות ביי
Top Hit Database Source	17	NT	1	Z	L	Ę	Ę	EST_HUMAN	EST_HUMAN	EST HUMAN	LZ	NT	۲۷	NT	NT	LN LN	LN	LN	N.	TN	LN	NT	NT	EST HUMAN	LN	۲	늏	LΝ	FZ	1:4	- !	Z	Z	Z
Top Hit Acession No.	5729928 NT	A DO 19330 1	T		0.0E+00(AB020717.1	7662237 NT	237	0.0E+00 AW 298134.1	-	0.0E+00 AA463659.1	0.0E+00 AB020710.1	7657468 NT	0.0E+00 AB037835.1	7662183 NT	4506718 NT	7857065 NT	T667085 NT	0.0E+00 AF195658.1		0.0E+00 7657468 NT	7657468 NT	4759011 NT	10181139 NT		0.0E+00 AF152496.1	4758199 NT	S78685.1	7710148 NT			00 AF069601.2	0.0E+00 AF069601.2	0.0E+00 AB001523.1	AB001523.1
Most Similar (Top) Hit BLAST E Value	0 0E+00	V 00-100	0.05-00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00/	0.0E+00	0.0E+00/	0.0E+00	0.0E+00		0.0E+00			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 S78685.1		00.0	0.0				
Expression Signal	1.53		1.81	1.1	1.1	22.21	22.21	4.35	4.35	1.08	1.14	3.31	0.82	6.4	4.29	1 08	1.08	0.71	2 88	2.3	2.3	1.24	1.01	1.01	1.97	1.14	10.39				1.23			0.97
ORF SEQ ID NO:	79000	١		29071	29072	29074				29105	L	L	L				8 8	L	20185							3 29198								
Exan SEQ ID NO:	10404	10450	16428	16430	16430	16432	16432	18445	16445	16467	16471	1	1	1	1_	. [_	L	L		1000+			L	1_	1	1	l	-			16576	18578		16580
Probe SEQ ID NO:	0200	20/25	3675	3877	3677	3679	3879	3692	3692	3714	3718	3721	3730	3742	3745	2754	3754		200	3802	2000	3808	3809	3812	3813	3814	3	3818	3819	3820	3824	3824	3829	3828

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	Top Hit Descriptor	Homo sapiens transient receptor potential channel 5 (TRPC5), mRNA	Homo saplens chromosome X open reading frame 5 (CXORF5) mRNA	Homo saplens chromosome X open reading frame 5 (CXORF5) mRNA	Human zinc finger protein ZNF134 mRNA, complete cds	Homo sapiens Intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1) mRNA	Homo sapiens familial mental retardation protein 2 (FWR2) gene, exon 11	Homo saplens SC35-Interacting protein 1 (SRRP129), mRNA	Homo saplens amphiphysin gene, partial cds	wk01f01.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:2411065 3' similar to TR: 043340 O43340 R28830_2, ;contains element PTR7 repetitive element;	Homo sapiens ribosomal protein S8 (RPS8), mRNA	DKFZp434N0413_r1 434 (synanym: htes3) Homo sapiens cDNA clone DKFZp434N0413 5'	Homo saplens AP1 gamma subunit binding protein 1 (AP1GBP1), mRNA	Homo sapiens AP1 gamma subunit binding protein 1 (AP1GBP1), mRNA	Homo sapiens glutamate receptor, metabotropic 3 (GRM3) mRNA	Homo sapiens melanoma antigen, family B, 1 (MAGEB1) mRNA	Homo sapiens HBP17 heparin-binding and FGF-binding protein gene, complete cds	Homo sapiens ryanodine receptor 3 (RYR3) mRNA	Homo sapiens zinc finger protein (KIAA0412) mRNA	RC3-HT0860-170800-011-a12 HT0860 Homo sapiens cDNA	MXRA5 Human matrix tissue expression library Homo sapiens cDNA clone incyte 1996726 similar to MXRA5. Matrix remodeling associated gene 5	MXRA5 Human matrix tissue expression library Homo sapiens cDNA clone Incyte 1996726 similar to MXRA5	Matrix remodeling associated gene 5	Homo sapiens F-box protein Fbt3b (FBL3B) mRNA, partial cds	UI-H-BI3-alh-g-07-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2736949 3'	601236968F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3608800 5'	PM3-LT0031-100100-003-h09 LT0031 Homo saplens cDNA	Homo saplens iduronate 2-sulfatase (Hunter syndrome) (IDS), transcript variant 1, mRNA	601193827F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3537774 5'	601193827F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3837774 5'	Human G2 protein mRNA, partial cds
	Top Hit Database Source	NT	TN	TN	NT	NT	FN	Ϊ́	Z Z	FZ	EST_HUMAN	N-	EST_HUMAN	TN	TN	N	ΝΤ	NT	LΝ	ΙN	EST_HUMAN	EST HUMAN		EST_HUMAN	LN	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	EST_HUMAN	NT
	Top Hit Acession No.	6912735 NT	4503178 NT	4503178 NT	U09412.1	AF114488.1	4826783 NT	1F01281	4759171 NT	AF099117.1	AI864727.1	4506742 NT	AL040338.1	6005887 NT	6005887 NT	4504138 NT	4505078 NT	AF149412.1	4506758 NT	4585842 NT	BF355295.1	AW888221.1		AW888221.1	AF129533.1	AW451306.1	BE378602.1	AW580740.1	5360215	BE264998.1	BE264998.1	U10991.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00
	Expression Signal	1.12	6.16	6.16	40.4	0.69	2.96	1.05	1.43	0.74	2.18	4.24	1.35	1.28	1.28	3.22	1.82	1.18	1.2	1.47	1.18	1.06		1.05	1.82	1	2.81	0.92	2.49	0.98	0.98	1.42
	ORF SEQ ID NO:			29223		29227		29232			29244	29248	29251			29269		59263	29273		29286	29288	İ	29289	29295	29298	29303	29311	29312	29313		29339
	Exan SEQ ID NO:	16582	16586	16586	16588	16589	16592	16595	16596	16598	16606	16609	16612	16617	16617	16819	16621	16625	16634	16638	16846	16648		16648	16654	16657	16662	16670	16672	16873	16673	16702
	Probe SEQ ID NO:	3831	3835	3835	3837	3838	3841	3844	3845	3847	3856	3859	3862	3867	3867	3869	3871	3875	3884	3888	3896	3898		3898	3904	3907	3912	3920	3922	3923	3923	3952

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Top Hit Descriptor	Human G2 protein mRNA, partial cds	Human G2 protein mRNA, partial cds	Homo sapiens cancer-testis antigen CT10 (CT10) gene, complete cds	Homo saniens cancer-testis antigen CT10 (CT10) gene, complete cds	Human MHC class II I Amphocyte antigen DPw4-beta-2 pseudogene, exon 2	1 lantar in the account of property of the second of the s	Home saplens critical action of the chomosome 20	Nove Italias yes company of commant HS24C084	Homo sapiens chromosome 21 segment HS21C068	Home septents concerned to beginned in the factor of alpha of (FEF1A4) mRNA	Homo sapiens eartalyout datistation across and a close MAAGE-224734 3' similar to TR: 060309 060309	Itabagus XI NCL COAF GOO name squares constructions in the KIAA0563 PROTEIN ;	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Human zinc finger protein ZNF133	1x781610 x1 NCI CGAP Lu24 Homo saplens cDNA clone IMAGE:2871694 3'	Characetus aethiops mRNA for ribosomal protein S4X, complete cds	Homo sapiens mRNA for UGA suppressor tRNA-associated antigenic protein (IRNA48 gene)	Homo sanians mRNA for KIAA0316 protein, partial cds	Homo sepiens chromosome 21 segment HS21C003	Home capitals media transcription factor 1-like (MYT1-l) mRNA, complete cds	Homo septens mRNA for rate-2 (rate dene)	Homo seniens mRNA for rane-2 (rane gene)	Tomo contone retinohlastoma-binding protein 4 (RBBP4) mRNA	Home contract retirobles from Ainding profess (RBBP4) mRNA	Home capiene G matella catalled recenter 21 (GPR21), mRNA	Hours cariers of program country of the country of	Troffic Squiets and CTD as antivetton protein-like (NGAP) mRNA	Hono saprens ras of rape activating process in a (IMPDH1) mRNA	Homo sapiens IMP (Inosine monophosphrate) derivatogenedo (Imp. 2017);	Nove human mixiva from critical lossements which has entired to the party of the pa	Homo sapiens Docked (Docked) micray, complete cos	Homo sapiens protein Kinase, A-linked (P-KNA) minna	Homo sapiens protein kinase, A-linked (P-RNA) miniva	Homo sapiens butyrophilin, subramily 3, merriber AZ (B.1N3AZ), minvry	Tono sapiens Christing process sanctipes in the same same same same same same same sam
Top Hit Database Source	Z	Į.	F	2 2	- E	Z	ŁV.	Ž	LN.	Į.	LN L	FST HUMAN	FZ	Į.	EST UIMAN	101	Z	1	- E	2	Z	2 1	- N	N.	Z	ĮN.	Z	LN.	Į.	ĻN.	Z	LN	L	NT NT	LN
Top Hit Acession No.]	T				ł	0.0E+00 AL163284.2	1163268.2	4503470 NT	0 0E 100 A 1857078 4	TAROTAR NT	1002100	000000000000000000000000000000000000000	0.0E+00 AW 338480.	0.0E+00 AB015610.1	0.0E+00/AJZ38617.1	0.0E+00 AB002314.2	0.0E+00 AL163203.2	00 AF036943.1	0.0E+00 AJZ77Z/6.1	00 AJ2772/6.1	5032026 NT	5032026 N	. 4885306 NT	AB00662		11419297 NT	0.0E+00 AL096857.1	0.0E+00 AF165527.1	4826947 NT	4826947 NT		4503854 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00 U10991.1	1 10001	0.05	0.05+00	0.0E+00	0.0E+00 MZ3910.1	0.0E+00 A	0.0E+00	0.0E+00 ≠	0.0E+00 /	0.0E+00	001110	00.00	0.0=+00	0.0=+00	0.0=+00.0	0.0E+00,	0.0=+00,	0.0E+00	0.0E+00.	0.0E±	1	0.0E±	}	-			0.0E+00	0.0E+00				0.0E+00		0.0E
Expression	1 42	,	1.42	4.69	4.69	4.39	5.74	1.35	3.22	1.71	27.98	7	2.10	2.97	2.65	0.95	6.33	3.72	1.32	1.04	1.18	2.65	2.65	6.29	6.29	4.7	5.98	1.11	6.87	1.94					1.08
ORF SEQ ID NO:	00000				28345							-					29401					29421				29442				L				29466	
SEQ ID	\perp	_				16715	ı	16724	1	16735	16747		⅃	_1	16752			16778		16791			16793	16799	١	16813	<u>L</u>	L	L	L	L		L	┸	ı
Probe SEQ ID NO:	0300	20AC	3952	3957	3957	3966	3968	3975	3979	3987	3899		4003	4005	4006	4013	4024	4033	4045	4048	4047	4048	4048	4054	4054	4069	4070	4073	4074	4075	4082	Ş Ş	800	4097	4099

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Top Hit Descriptor (GABPA), mRNA	Homo sapiens GA-binding protein transcription recent in the contract of KIAA0895 protein, partial cds	Mundado4 x1 NCI CGAP GC6 Hamo sapiens cDNA clone IMAGE:2515975 3	wn04604.x1 NG, CGAP, GC6 Homo sapiens cDNA clone IMAGE:2515975 3	MR1-HT0707-100500-001-602 HT0707 Hamo sapiens cDNA	MR1-HT0707-100500-001-a02 HT0707 Homo sapiens CDNA	801120778F1 NIH_MGC_20 Home sapiens cDNA clone IMAGE: 2967690 3	Homo sapiens mRNA for KIAA1125 protein, partial cds	Homo sapiens mRNA for KIAA1125 protein, par usa wa	Homo sapiens nuclear receptor coecityator 3 (NCOA3), minya Tana Mange 2900095 3' similar to SW:THI2_BOVIN	bash04.x1 NIH_MGC_10 Home sapiens curva dare into curva and into curva mitrochondrial THIOREDOXIN PRECURSOR;	UB3108 WILLOW 1000 1000 1000 1000 1000 1000 1000 10	Homo sapiens hypothetical protein FLJ10498 (FLJ10498), mRNA	Homo saplens hypothetical protein FLJ10498 (FLJ10498), mRNA	Homo spalens balveystic lidney disease (polycystin) and REJ (sperm receptor for egg jelly, sea urchin)	homolog)-like (PKDREJ) mRNA homo septens cDNA clone IMAGE:743197 3' similar to contains Alu	Zu68h07.st Sogres, lesus, 1411 i tonico de peditivo element ;	repolutive destress, contains All 2016	repetitive element contains element MER35 repetitive element ; Home seplens chromosome 21 segment HS21C103	Human and locarotein B-100 mRNA, complete cds	PM2-DT0023-080300-004-a08 DT0023 Homo saplens cDNA	Homo sapiens myelodysplasia syndrome 1 (MDS1) mRNA	Homo sapiens myelodysplasia syndrome 1 (MDS1) mRNA	Homo sapiens F-box protein Fb/4 (FBL4) mRNA, partial cds	qd23f06,x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens curva clure invocation	$\neg \tau$	Т	Home sabiens protein kinase C, nu (PRKCN), mRNA	Homo septiens protein kinase C, nu (PRKCN), mRNA	Human G2 protein mRNA, partial cds	
Top Hit Database Source	Į.	NI FOT UNAN	EST FINANI	EST HUMAN	EST HIMAN	EST HUMAN	E	LN.	LN		EST HUMAN	אויים ביי	Z	Ž	님		EST HUMAN	EST_HUMAN	Z I	EST HIMAN	-1	TN P	LN LN		EST_HUMAN	N	6 NT	4 .	NAT NAT	
Top Hit Acesslon No.	3854		Al982597.1	A1982597.1				AB032951.1	5729726 NT		AW675599.1	AW408788.1	8922466 NI	8922466 N	5174632 NT		AA401438.1	AA401438.1	AL163303.2	J02610.1	0.0E+00 AW \$30009.1		A E 4 7 4 50	0.0E+00/AF1/4090.1	0.0E+00 AI189844.1	0.0E+00 U14520.1	4505646 N1		6563384 N	00101000
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00 AI		0.0E+00 A		0.0E+00 B		V 00730	0.0F+00		0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 J02610.1						L			0.0E+	0.00
Expression Signal	1.08	0.89	4.93	4.93	0.82	0.82	2,34	0.99	0.89	27.7	5.52	1.14	1.26	1.26	2.8		8.97	8.97	1.01				٥	4.7	2.52	4.32				1.58
ORF SEQ ID NO:	29470	29476	29482		29485	29486				29490		29511		L			29537	29538		29606				4 29636		0 6	29650			36 29664
Exan SEQ ID NO:	16842	16850	16855	16855	16857	L			1	16870	16877	l	1	1_	1	1	16909	16909		1_	16996	1 17002		3 17004		1/010	┸	1_	L	Ы
Probe SEQ ID NO:	4000	4107	4112	4112	4115	4115	4120	4126	4126	4128	4135	4140	4142	4142		4151	4169	4469	4205	4240	425	4261	4261	4263	!	4270	4213	300	4291	4297

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Top Hit Descriptor	Human G2 protein mRNA, partial cds	Hamo sapiens COMPLEMENT COMPONENT C14 RECEPTOR (C1QR), mKNA	Homo sepiens gap junction protein connextn-36 (CX36) gene, complete cds	Home contains placema mambrane calcium ATPase isoform 1 (ATP281) gene, alternative splice products,	nonto espesas pressina mentra espesas partial cds	H.saplens H2B/h gene	H.saplens H2B/h gene		H. sapiens H4/d gene for H4 histone	H.sapiens H4/d gene for H4 nistone	Homo sapiens KIAA0390 gene product (NIAA0390), IIINAA	Homo saplens KIAA0390 gene product (KIAA0390), mKNA	Homo sapiens caudal type homeo box transcription factor 4 (CDA4), minner	Homo sapiens Xq pseudoautosomal region; segment 2/2	Homo saplens myosin regulatory light chain interacting protein (vinity, interacting protein (vinity), interacting protein (vin	Homo sapiens membrane-bound aminopeptidase P (ANPERZ) garie, Compress cus	Homo sapiens ACTN2 gene for alpha-Actinin 2, expn 10	Homo sapiens ACTNZ gene for alphe-Actinin Z, excn 10	Homo sapiens HPS1 gene, intron 5	П	\neg	Human endogenous retrovirus nervy no			wc66b02.x1 NCI_CGAP_Pr28 Homo sapiens cDNA clone inviaGE_2322503.3 similar v community minimum in TRS penetitive element:	Т	Т	7	United applies mirror or programmer and a second containing protein (ORF1)	House anniens G protein-criminal recentor 50 (GPR50) mRNA	Homo septents serine-threoning protein kinase (MNBH) mRNA, complete cds	
Top Hit Database Source	LN	LZ.	17		LN.	S	-LZ	EST_HUMAN	LN	N	NT	۲N	ΝΤ	LN.	LN	LN	NT	N	TN	EST_HUMAN	EST_HUMAN	Z	EST_HUMAN	NT	ECT LIMAN	ESI TOWN	NAAN LI HAAN	Caron 1 22	- !	Z	Z	
Top Hit Acessian No.	001110991.1	2281	0.05 +00	T			0.0E+00 Z80780.1	AW166933.1	+00 X60483.1	0.0E+00 X60483.1	7662091 NT	7662091 NT	4885126 NT	AJ27173	7019456 NT	-00 AF195953.1	-00 AJ249765.1	-00 AJ249765.1	00 AF200629.1	-00 T10233.1	-00 T10233.1	HOD M14123.1	+00 AW084964.1	8051619 NT	1000000	100 Albeober 1	F00 ALTOSZU7.2	+00 AW 3613/0.1	+00 AJ278120.1	AJ27812	4758467 NI	+00/AF-108830.1
Most Similar (Top) Hit BLAST E Value	001100	100	0.00	0.00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.00+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E	0.0E	1	G.O.	0.011400	0.0	0.0E	9.0	9.0	0.0E
Expression Signal	4 60	3	10.0	1.16	4.17	2 88	2 88	0.94	1.42	1.42	8.62	8.62	12.59	1.14	0.98	6.9	1.25	1.25	1.81			0.73		1.67								2.88
ORF SEQ ID NO:	1	28000	١			00/02	L		29717	29718		Ŀ					29778			29814			20828			5 29831				3 29843		9 29846
SEQ ID	OG GE,	000/	-1	17068	4707B	17007	77080	ł	1	1_	L		丄	1	L	1_	L	1	_L_	L	١.		l	1	L			17210	17216	17216	1 3	4 17219
Probe SEQ ID NO:		187 ₄	4307	4327	4937	3	2	5	4348	4348	1253	4252	AS CA	4367	740	AAA	777		4414	4453	4453	4456		4468		4470	4473	4475	4481	4481	4483	4484

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Table 4
Single Exon Probes Expressed in Brain

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	Top Hit Descriptor	Homo sapiens ATP-sensitive inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, exon	Homo sapiens pyrin (MEFV) gene, complete cds	Homo sapiens pyrin (MEFV) gene, camplete cds	Homo saplens zinc finger protein 195 (ZNF195), mRNA	Homo saplens syncytin precursar, mRNA, camplete cds	Homo sapiens protocadherin gamma C3 (PCDH-gamma-C3) mRNA, complete cds	Homo sapiens zinc finger protein 211 (ZNF211), mRNA	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo sapiens low density lipopratein receptor-related protein 6 (LRP6) mRNA, and translated products	Homo sapiens chondroitin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA	Homo sapiens calcium/calmodulin-dependent protein kinase IV (CAMK4) mRNA	Homo sapiens iduronate sulphates sulphatase (IDS) gene, complete cds	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens PTEN (PTEN) gene, exons 3 through 5	Homo saplens mRNA for G7c protein (G7c gene located in the class III region of the major histocompatibility complex)	Homo saplens mRNA for G7c protein (G7c gene located in the class III region of the major histocompatibility	complex)	zp18g08.s1 Stratagene fetal retina 837202 Homo sapiens cDNA clone IMAGE:609854 3'	Homo sapiens odz (odd Oz/ten-m, Drosophila) homolog 1 (ODZ1), mRNA	Homo saplens chromosome 21 segment HS21C084	y492b01.s1 Soares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:231721 3'	y92b01.s1 Soares_pineal_gland_N3HPG Homo sapiens cDNA clone IMAGE:231721 3'	Homo sapiens cyclophilin-related protein (NKTR) gene, complete cds	Homo saplans chromosome 21 segment HS21C100	Homo saplens gene for natriuretic protein, partial cds	Homo sapiens keratin 18 (KRT18) mRNA	Homo saplens keratin 18 (KRT18) mRNA	Homo sepiens inwardly-rectifying potassium channel Kirz.1 (KCNJ2) gene, exon 2 and complete cds	Homo sapiens inwardiy-rectifying potassium channel Kir2.1 (KCNJ2) gene, exon 2 and complete cds	
201 1110000 216	Top Hit Database Source	NT	LN	NT	٦	NT	TN	TN	LN	LN	LN	LN LN	L	N.	LN LN	N _T	ĘN			T_HUMAN	NT		EST_HUMAN	EST_HUMAN	NT		NT	LN	NT	NT	TN	
	Top Hit Acession No.	S78684.1	AF111163.1	AF111163.1	TN 5765009	AF208161.1	AF152337.1	5454175 NT	4503470 NT	4505016 NT	4503098 NT	4502556 NT	35485.1	7662091 NT	7662091 NT	AF143314.1	AJ245418.1		AJ245418.1	AA174072.1	- 7657410 NT	AL163284.2	H92741.1	H92741.1	AF184110.1	AL163300.2	AB037521.1	4557887 NT	4557887 NT	AF153819.1	AF153819.1	
	Most Similar (Top) Hit BLAST E Value		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_		0.0E+00	0.0E+00		0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	
	Expression Signal	1.26	1.06	1.06	2.56	6.16	4.31	1.32	15.47	0.79	1.61	2.03	2.38	12.72	12.72	96.0	10.33		10.33	1.68	1.46	3.16	1.04	1.04	2.8	4.94	1.66	1.53	1.53	1.52	1.52	
	ORF SEQ ID NO:		65862	29860	29870		29882	29886	29894	29900	29903	29908		29910	29911	29925	29928		29929				29947	29948	29949	28950		29958	29959	29960	29961	
	Exan SEQ (D NO:	17229	17230	17230	17882	17242	17247	17250	17260	17267	17271	17275	17279	17281	17281	17298	17301		17301	17316	17318	17320	17321	17321	17322	17323	17324	17331	17331	17332	17332	
	Probe SEQ ID NO:	4493	4494	4494	4502	4507	4512	4515	4525	4532	4536	45 545	4544	4546	4546	4563	4566		4566	4581	4583	4585	4586	4586	4587	4588	4589	4596	4596	4597	4697	1

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Single Exon Probes Expressed in Brain

Human proto-oncogene tyrosine-protein kinase (ABL) gene, exon 1a and exons 2-10, complete cds SCN1A=brain type I sodium channel alpha-subunit {IIIS6 transmembrane region} [human, placenta, Genomic SCN1A=brain type I sodium channel alpha-subunit {IIIS6 transmembrane region} [human, placarta, Genomio Homo sapiens truncated tenascin XB (TNXB) gene, partial cds and TNXA gene recombination breakpoint Mus musculus neurexophilin 1 (Nxph1) gene, large exon and 3' end of the Intron, and partial cds ya83g04.r2 Stratagene fetal spleen (#937205) Homo saplens cDNA clone IMAGE:68310 6' 801158935F1 NIH_MGC_21 Homo saplens cDNA olone IMAGE:3505521 5' Homo sapiens bromodomain adjacent to zinc finger domain, 28 (BAZ2B), mRNA Homo sapiens alpha-3 type IX collagen (COL9A3) gene, promoter region, and exons 1-26 Homo sapiens proteinx0008 (AD013), mRNA ya83g04.r2 Stratagene fetal spleen (#937205) Homo sapiens oDNA clone IMAGE:68310 Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds Homo saplens bromodomain adjacent to zinc finger domain, 2B (BAZ2B), mRNA Human endogenous retrovirus type K (HERV-K), gag, pol and env genes zv86b07.s1 Soares_NHHMPu_S1 Homo sapiens cDNA clone IMAGE:767606 3' Homo sapiens butyrophilin, subfamily 2, member A2 (BTN2A2), mRNA Homo saplens cyclophilin-related protein (NKTR) gene, complete cds Homo saplens KIAA0563 gene produot (KIAA0563), mRNA Mus musculus E-cadherin binding protein E7 mRNA, complete cds Top Hit Desoriptor Homo sapiens butyrophilin, subfamily 2, member A2 (BTN2A2) Human CYP2D7AP pseudogene for cytochrome P450 2D6 Homo sapiens chromosome 21 segment HS21C078 Homo sapiens mRNA for KIAA1399 protein, partial cds Homo sapiens mRNA for KIAA1047 protein, partiel cds Homo sapiens mRNA for KIAA1047 protein, partial cds Homo saplens mRNA for KIAA1399 protein, partial ods Homo sapiens chromosome 21 segment HS21C078 Human AHNAK nucleoprotein mRNA, 6' enc 1556 nt] 1556 nt] region EST_HUMAN NT HUMAN **EST_HUMAN** HUMAN Top Hit Database Source EST 눋 눋 눌 E 巨巨 Ę 7304922 NT z 눋 7662181 NT 6453812 N 7019320 7304922 Top Hit Acession 0.0E+00 AL163278.2 0.0E+00 AB037820.1 0.0E+00 AB037820.1 0.0E+00 AB028970.1 0.0E+00 Y18890.1 0.0E+00 AA418246.1 0.0E+00 AF167441.1 0.0E+00 AB028970.1 AF026801.1 0.0E+00 AF086641.1 0.0E+00 AL163278.2 AF184110. 0.0E+00 BE278730. 0.0E+00 U56651.1 0.0E+00 M80902.1 0.0E+00 M69197.1 ջ 0.0E+00 T56945.1 0.0E+00 BE278730 T56945.1 0.0E+00 M74099.1 0.0E+00 U07583.1 S71446.1 0.0E+00 S71446.1 X58467 0.0E+00/ 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLASTE Most Similar 0.84 1.45 1.05 .05 1.06 0.82 0.95 0.97 0.97 1.22 1.22 5.25 1.08 2.72 1.84 1.31 Expression Signal 30073 30076 30065 30066 30049 ORF SEQ ID NO: 29971 29979 30001 30002 30005 30006 25593 30038 30040 30050 30051 29970 29987 17415 17415 17435 17414 17441 17340 17333 17340 17359 17370 12950 17399 SEQ (D) 17364 17370 17352 ğ 4709 4639 4660 4668 4668 4671 4692 4701 4805 805 805 4630 4631 4635 4681 4701 4629 4629 4635 4636 4681 4611 4824 4630 4636 4673 88 SEQ ID 4598 ë

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	Top Hit Descriptor	Homo sertiens proteinx0008 (AD013), mRNA	UI.H-BI3-alw-c-04-0-UI.s1 NOI_CGAP_Sub5 Homo saplens cDNA clone IMMCE. 2 3227-7	Hamo sapiens HSPC024-Iso mRNA, complete cds	Human connexin 43 processed pseudogene	Homo sapiens glutathione S-transferase theta 2 (GST12) and glutathione S-transferase theta 2	genes, complete cds M feerinalaris mRNA for metalloprotease-like, disintegrin-like protein, IVa	Homo sapiens Williams-Beuren syndrome deletion transcript 9 (W BSCR9) mrdvA, complete cus Homo sapiens Williams-Beuren syndrome deletion transcript 9 (W BSCR9)	Mus musculus zinc finger transcription factor Kalso minky, compress zinc finger transcription factor Kalso minky,	Homo sapiens fragile X mental retardation 2 (FMKZ) minus	Homo sapiens actin, alpha, cardiac muscie (Actic), misson	ZING FINGER PROTEIN 132	Homo sapiens hypothetical protein FLJ20073 (FLJ20073), mixivo	Homo sapiens KIAA0187 gene product (KIAA0187), mKNA	Human Tor-C-detta gene, exons 1-4, Tor-V-delta gene, exons 1-2, 1-cal receptua apria (15, 15, 15)	J61 segments; and Tcr-C-alpha gene, exons 1-4	Human Tor-C-delta gene, exons 1-4; Tor-V-delta gene, exons 1-2;	J61 segments; and Tor-C-alpha gene, exons 1-4 H sabiens MeCP-2 gene	H.saplens MeCP-2 gene	Homo sablens chromosome 21 segment HS21C080	Home segiens TATA box binding protein (TBP)-associated factor, RNA polymerase II, I, cond. (1787)	mRNA	H.sapiens MICA gene	Homo sapiens zinc finger protein (NIAANU412) in the	Homo sapiens mRNA for KIAA0633 protein, partial cus	Mus musculus zinc finger protein interacting with it protein (Cir.), MRNA	Homo sapiens meningloma expressed angger o (varied or promise)	Homo sapiens desmoplatin (DPI, DPII) (DSF) minuto	Homo sapiens zino-finger DNA-binding protein (110 in 100 100 100 100 100 100 100 100 100 10	Homo saplens MHC class 1 region	Hano saplens opiold receptor, delta 1 (OPRD1) mrvns	Homo sapiens spilce variant AKAP350 mRNA, parted cus	Homo sapiens COL4A6 gene for a6(IV) collagen, exon 44 and parted con	
Single Exoli r lobe	Top Hit Database Source		T LI MAAN	Т	FIN		LZ!	Z	LZ.	NT.	TZ.	SWISSPROT	Į.	Į.	2	LZ		F	2 12		Z	F2	LZ	NT	L _N	8 NT	TNIC	TN	LN 9	LZ	LN	LZ	Į.	
Buis	op Hit Acession No.		222	W444637.1		M65189.1	5.1	X87205.1	AF007446 1	4503766INT	ARREDAR NT	250240	TN OBOSCOO	9923000	INIB/BLOO/	MOADR4 4	10001.1	M94081.1	X94628.1	X94628.1	AL163280.2	TULOSOS	VO2841 4	AFREGAZINT	0.0E+00	6677848 NT				ACOECOE 4	0.0E+00 A-033000.	A 1700474 4	0.0E+00 AFUST 11.1	
		Value	1	~ 1 7		0.0E+00		_		0.01	0.05	0.0E+00		0.0E+00	0.0E+00		0.0E+00.	0.0E+00		0.0E+00	0.0E+00		0.01+00	0.01	0.00	0.05+00	0.0	1				١	1	
	Expression (Signal		0.84	1.88	1.82	2.28	2.79	2.02	1.11	1.96	3.01	13.57	1.04	5.7	76.0		1.84	1.84	1.44	1.44	2.96					1.81						3.47		1.07
	ORF SEQ E	_	30077	30103				30147	30149				30154		30163	_	30164	30165				L						1			72 30196	4		37 30210
	0		17444	17467	17472	17482	17521	17525	17527	17528	_	17531	<u>L</u>	1	١_		17541	17541	L	L	L	L	3 17554		<u> </u>	3 17564	17565		17567	17569	17572	47574	45 17575	58 17587
		ö	4712	4735	4740	4750	7,700	4794	4798	4797	4798	88 88	4801	4805	4809		4810	7870	C 184	4612	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		4823	4830	4832	4833	4834	4835	4836	4838	4842	4844	4845	4858

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wo	01/57275		┯-	- ī	- -	_	_		_	_	_	_	_		_	_		_BB_	- H	****]	P <u>C</u>	T/U	SO,	1/0	066	<u> </u>
Table 4 Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens farnesyl diphosphate synthase (farnesyl pyrophosphate synthetase, dimethylallytranstransferase, geranytransfransferase) (FDPS) mRNA	Homo saplens sialytransferase 8 (alpha-N-acetylneuraminate: alpha-2,8-sialytransferase, GD3 synthase)	Homo saplens mRNA for KIAA0287 mens mental such	Homo sapiens mRNA for KIAA0287 gene, partial cds	Homo sepiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	Homo sapiens chromosome 24 sommet UC240004	U-H-Bl3-glwf-02-0-U1 s1 NCT CGAP Subs Home content and state 1440 Frances of	Homo saplens bypothetical protein El 144400 /El 144400 BMA	Homo saplens beaded filament structural protein 1 filensin (RESDA) mibria	Human ribosomal profesion 24 mRNA complete of 1, minuted	601303729F1 NIH MGC 21 Homo sepiens CDNA close IMAGE-3838449 ET	Homo saplens desmonakin (DPI DPII) (DSP) mBNA	Homo saciens KIAA0952 protein (KIAA0952) mena	Homo sapiens mRNA for KIAA1043 motion particled	Homo seplens hyndrhetical notein El (2017) (El 19042) - BNA	Homo saplens hypothetical protein FL J20477 (FL J20477), mRNA	no14g08.s1 NCI CGAP Phe1 Hamo sapiens cDNA clone IMAGE:1100704 3' similar to TR:E239140 E239140 SPALT PROTEIN :	no14g08.s1 NCI_CGAP_Phe1 Homo saplens cDNA clone IMAGE:1100704 3' similar to TR:E239140 E239140 SPALT PROTEIN :	no14g08.s1 NCI_CGAP_Phe1 Homo saplens cDNA clone IMAGE:1100704 3' similar to TR:E239140 E239140 SPALT PROTEIN :	Homo saplens HSPC/14 mRNA complete cds	Homo sapiens HSPC114 mRNA complete cde	Homo sapiens DNA mismatch repair protein (MLH3) gene complete cds	H.sapiens mRNA similar to D29763 mouse mRNA for seizure-related gene product 6. Shares domeins with	Divir s, Tolido, Sushi repeat proteins Homo saplane E8 40 Historial	Mus musculus mRNA for extrall short animal version for the state of th	Mus musculus mRNA for aczonin, short spliced variant fact gene)	Homo sapiens chromosome 21 segment HS21C009
Igle Exon Pro	Top Hit Database Source	ΓN	L _Z	NT.	LN.	ΤN	LN	EST HUMAN	Ę	F	Ľ	EST HUMAN	NT	N.	NT	Į.	Ļ	EST_HUMAN	EST_HUMAN	HUMAN	T		N	- FN				± L
Sir	Top Hit Acession No.	4503684 NT	4506952 NT	AB006625.1	0.0E+00 AB006825.1	0.0E+00 AB026898.1	VL163284.2	W 452728.1	8922926	4502398	114967.1	E408863.1	4758199 NT	7662401 NT	28966.1	8923441 NT	8923441 NT	0.0E+00 AA601246.1					95658.1			Γ		
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00/	0.0E+00 AL	0.0E+00 AW	0.0E+00	0.0E+00	0.0E+00 U14	0.0E+00 BE	0.0E+00	0.0E+00	0.0E+00 AB0	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 AA801246.1	0.0E+00 AA601246.1	0.0E+00 AF161483.1	0.0E+00 AF161463.1	0.0E+00 AF1	0.0F+00 AI 050253 1	0.0E+00 AF016705.1	0.0E+00 Y191	0.0E+00 Y191	0.0E+00 AL163209.2
	Expression Signal	1.68	1.03	3.09	3.09	0.95	1.45	1.02	1.61	1.09	4.69	2.95	3.18	1.15	1.01	2.34	2.34	0.81	0.81	0.81	1.11	1.11	0.71	0.84	1.63	1.5	1.5	1.28
	ORF SEQ ID NO:	30212			30225	30232	30243	30248	30251	30255			30268	30271	30274	30283	30284	30291	30292	30293	30295	30296	25661		30312	30313	30314	
	Exon SEQ ID NO:	17589			17602				_		- 1	17652	17656	17661	17666	17674	17674	17683	17683	17683	17686	17686	13018	17699	17708	17709	17709	17717
ł	Probe SEQ ID NO:	4860	4865	4875	4875	4885	4899	4906	4909	4912	4915	4824	4928	4833	4938	4947	4847	4958	4958	4958	4961	4961	4973	4976	4985	4986	4986	4994

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Single EXUITIONES Expressed III Didiri	Top Hit Descriptor	Homo sapiens gammma-cytoplasmic actin (ACTGP3) pseudogene	zn03g10.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:546402 5	zn03g10.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:546402 5	Homo sapiens titin (TTN) mRNA	Homo sapiens titin (TTN) mRNA	Bacillus amyloliquefaciens sacB gene for levansucrase (EC 2.4.1.10)	Homo sapiens vascular endothelial cadherin 2 mRNA, complete cds	Homo sapiens vascular endothelial cadherin 2 mRNA, complete cds	Homo saplens ecotropic viral integration site 2A (EVI2A), mRNA	Homo sapiens ecotropic viral integration site 2A (EVI2A), mRNA	Homo sapiens chromosome 21 segment HS21C081	Homo sapiens HEF like Protein (HEFL), mRNA	Homo saplens PR domain containing 1, with ZNF domain (PRDM1) mRNA	Homo sapiens putative GPR37 gene, exon 2	Homo sapiens putative GPR37 gene, exon 2	Human endogenous retrovirus-K, LTR US and gag gene	Homo sapiens potassium inwardly-rectifying channel, subfamily J, member 16 (KCNJ16), mRNA	Homo saplens potassium inwardly-rectifying channel, subfamily J, member 16 (KCNJ16), mRNA	Homo sapiens 4F2 light chain (LOC51597), mRNA	Homo sapiens 4F2 light chain (LOC51597), mRNA	Homo saplens defeted in bladder cancer chromosome region candidate 1 (DBCCR1), mRNA	Homo sapiens mRNA for KJAA0559 protein, partial cds	Mus musculus mRNA for leucine-rich repeat protein, partial cds	Rattus norvegicus multidomain presynaptic cytomatrix protein Piccolo mRNA, complete cds, long spilce variant	Raftus norvegicus multidomain presynaptic cytomatrix protein Piccolo mRNA, complete cds, long splice	variant	Homo sapiens toll-like receptor 7 (TLR7) mRNA, complete cds	Homo saplens microtubule-associated protein 2 (MAP2) mRNA		tate 2A (GRIN2A) mRNA	s cDNA	artial cds	Homo sapiens gababr1 receptor gene, exon 6
פופ באטוו רוטב	Top Hit Database Source	NT	EST_HUMAN	EST_HUMAN	NT	TN	NT	NT	NT	IN	1N	NT	NT	NT	ΙN	TN	NT	TN	NT	LN	TN	NT	NT	NT	ΙN		NT	IN	NT	NT		EST_HUMAN	Ę	Ė
110	Top Hit Acession No.	D50657.1	AA084272.1	AA084272.1	4507720 NT	4507720 NT	X52988.1	AF240635.1	AF240635.1	7657074 NT	7657074 NT	AL163281.2	11421001 NT	4557362 NT	Y12477.1	Y12477.1	Y08032.1	8923822 NT	8923822 NT	7706245 NT	7706245 NT	7657008 NT	AB011131.1	D49802.1	AF227534.1		AF227534.1	AF245702.1	4505096 NT	6006002 NT	6006002 NT	AW955819.1	AB040946.1	AJ010179.1
	Most Similar (Top) Hit BLAST E Vatue	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00/
	Expression Signal	26.03	76.0	76.0	0.95	0.95	3	1.04	1.04	1.55	1.55	1.11	14.05	1.03	2.75	2.75	1.07	1.01	1.01	97.0	0.76	2.69	2.05	1.23	1.14		1.88	66.0	6.53	1.5	1.5	1.6	1.31	1.12
	ORF SEQ ID NO:		30326	30327	29553	29554	30359	30375	30376	08606	30381	28808	88606	30389	30391	30392	30394	30414	30415	30417	30418	30423		30444	30445		30446	30447	30449	30450	30451			
	Exan SEQ ID NO:	17720	17724	17724	16924	16924	17747	17761	17761		17764		17769	12221	17775	17775	17777	17798	17798	17800	17800	17807	17816	17827	17828	ł	17829	17830	17833	ŀ	17834		_	17844
	Probe SEQ ID NO:	4997	5001	5001	5012	5012	5026	5042	5042	5045	5045	5049	2050	5052	2056	9909	8909	6209	5079	5081	5081	2088	2009	5109	5110		5111	5112	5115	5116	5116	5117	5119	5128

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wo	01/57275	7		7	.		_	T	-	Т	T					_		_,	-	<u> </u>		P .	وین آ	F	<u>S</u>	Γ/(→	ĴS(01/	00	66	7 * *
Table 4 Single Exon Probes Expressed in Brain	Top Hit Descriptor	Hamp sapiens mRNA for Nierlanden Angemehr Bratele 4 III.	Homo saplens mBNA for no mode of a large of the contract of th	Homo sapiens mRNA for KIAA1117 profela Add	Human sodium channel mRNA	Homo sabiens ring finger profeln 15 (RNE4R) mBNA	Homo sapiens aconitase (ACO2) gene. nuclear gene encoding milrohanddol ametric accentate	Homo sapiens keratin 12 (KRT12) gene, complete cds	Homo saplens keratin 12 (KRT12) gene, complete cds	Wp08g08.x1 NCI_CGAP_Kid12 Homo saplens cDNA clone IMAGE:2464094 3'	Homo sapiens protocadherin alpha 13 (PCDHA13), mRNA	Homo seriens notward Aldred III 1	Home seriens pulycysus ranged alsease-like 2 protein (PKDL2) mRNA, complete cds	H. Saplens immunodolphilin heavy chair and watch and watch.	H. sabians immunociotalin beaus chain sees, vertable region	710c08.x1 NCI CGAP CL1.1 Homo septems CDM & close MA CE (2004/25) 21	https://www.misser.com/septembers/misser.com	601589422F1 NIH MGC 7 Homo seniens cDNA class MAACE 2013004 51	601589422F1 NIH MGC 7 Homp sablens CDNA Clone IMAGE 3043804 5	4d04a04.x1 Soares_placenta_gto9weeks_ZNbHP8to9W Homo saplens cDNA clone IMAGE:1722702.3'	Homo septems excluded be and the Part of the Control of the Contro	Homo sepiens Sp4 francription furth (SBA)BNA	602118928F1 NIH MGC 56 Home series - DNA class MA CELESCAL 2	AU134406 OVARC1 Home services CONA Clone Invite Services	AU134406 OVARC1 Homo seniens CDNA clans OVARC1001884 5	601061489F1 NIH MGC 10 Home sanians cDNA close IMAGE:247222 F	601105891F1 NIH MGC 15 Home seniers CDNA clore MACE: 2089342 E	302071372F1 NCI	802071372F1 NCI_CGAP_Bm64 Homo saplens cDNA clone IMAGE-4214372 8'	Homo saplens Bloom syndrome (BLM) mRNA	Homo sapiens mRNA for KIAA0466 protein, partial cols
ğle Exon Pro	Top Hit Database Source	Ľ.	L	Z	LN L	NT	N	NT	ا ا	EST HOMAN	ŀ	NEW LONG	12	Z-L	LN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HIMAN	LN LN	L	EST HUMAN	11		EST HUMAN	EST HUMAN	Г	T_HUMAN		Į
Sin	Top Hit Acession No.	0.0E+00 AB027013.1	0.0E+00 AB035356.1	0.0E+00 AB029040.1	0.0E+00 M91803.1	5454013	93093.1	37286.1		0.0E+00 Al934954.1 ES	9/2002/B	82034.1	82034.1			2		0.0E+00 BE794412.1	l			421038	35962.1	34406.1			Γ		5328.1	4557364	7935.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AFC	0.0E+00 AF	0.0E+00 AF1	0.00+00.0	0.0E+00	0.0E+00 AF	0.0E+00 AF	0.0E+00	0.0E+00 X56163.1	0.0E+00 BE6	0.0E+00 E	0.0E+00 E	0.0E+00 E	0.0E+00 A1189142.1	0.0E+00 MZ9908.1	0.0E+00	0.0E+00 BF66	0.0E+00 A	0.0E+00 AU134406.1	0.0E+00 BE538857.1	0.0E+00 B	0.0E+00 BF5	0.0E+00 BF52	0.0E+00	0.0E+00 A
	Expression Signal	5.57	1.19	1.18	1.08	1.37	3.44	2.52	2.52	77.1	3.81	3	6	1.66	1.66	6.56	1.75	1.93	1.93	0.81	6.17	4.68	7.18	0.73	0.73	-	1.07	1.69	1.69	1.82	0.91
	ORF SEQ ID NO:	30473	30477	30481		30484		30495	30496	30628	30639	30643	30644	30652	30653	30763	30764	30765	30766	30767	30770	. 30780		30791	30792	30799	30826	30831	30832	32513	30858
	Exen SEQ ID NO:					- 1	- 1	- 1	18001	1	1		1	18027	18027	18104	18105	18108	18108	18107	18111	25066	18132	18133	18133	18138	18147	18151	18151	19491	18171
	Probe SEQ (D NO:	6139	6150	5151	5164	5155	5162	01/2	5192	5195	5209	5213	6213	5220	5220	6239	5300	6301	889	5302	5308	5319	6329	2330	2330	5335	5344	5348	5348	5367	6370

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Most Similar Top Hit Acession Ontabase ID ID NO: Signal Volve		30859 0.91 0.0E+00 AB007935.1 ivi	30863 4.85 0.0E+00 AF257737.1 NI	30864 4.85 0.0E+00 AF25/73/.1	30878 1.06 0.0E+00 D28535.1 INI	30879 1.06 0.0E+00 D26535.1 INI	30906 1.88 0.0E+00 11420819 N.1	30912 0.81 0.0E+00/238133.1 N1	30936 0.78 0.0E+00 D81664.1 EST_HUMAN	20027 0 78 0.0E+00 D61564.1 EST_HUMAN	30931 2.55 0.0E+00 BF529931.1 EST HUMAN	30942 2.65 0.0E+00 BF\$28931.1 EST_HUMAN	30948 2.92 0.0E+00 BF313139.1 EST_HUMAN	31132 4.37 0.0E+00 11434392 NI	31171 1.16 0.0E+00 BE260777.1 ESI HUMAN	3.51 0.0E+00 AW80/310.1	31196 2.33 0.0E+00 BE292889.1 E31_FCWAN	2.33 0.0E+00 BE282883.1 EO:	31219 1.51 0.0E+00 114203191VI	31220 1.51 0.0E+00 114200.9 NT	4.30 0.0E+00 AF0640E4 4 INT	31229 4.35 0.0E+00 A 1224839.1 NT	31239 2.30 C.30 NT		1 EST HUMAN	31277 6.98 0.0E+00 M85719.1 EST HUMAN	31286 4.83 0.0E+00 AW 403472.1	31296 1.25 0.0E+00 226269.1 N.	0.0E+00 AW3618//.1	31307 1.94 0.0E+00 AW3618/7.1 EST DIMAN	31308 1.94 0.0E+00 AW361877.1 EST_DUMPAY	
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Exon D SEO ID		70 18171	L	<u> </u>		L	1_	L	5426 18225	١	5426 18225	5429 10220	L	1_		5486 18285	5500 18298	乚	L	Ш	6528 1832			5535 1833	25 1836 1836	L		L	<u> </u>			Ш
Probe SEQ ID	<u>.</u>	6370	6374	5374	5387	5387	5401	R	7		الم	ည် မြိ	5 1	7	Č.	ů	, and	9	۳	ا ا	Φ	(°)	ųΣ	3)								

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	Top Hit Descriptor	Homo saplens mRNA for KIAA1641 protein, partial ods	Homo sabiens KVLQT1 gene	Homo sapiens KVLQT1 gene	HA2981 Human fetal liver oDNA library Homo saplens cDNA	Homo saplens protocadherin beta 2 (PCDHB2), mRNA	601584032F1 NIH_MGC_7 Homo saplens oDNA clone IMAGE: 3838001 9	Homo saplens amiloride-sensitive cation channel 1, neurona (degenerin) (NOON).	601346141F1 NIH_MGC_8 Homo saplens cDNA digna liwa on 1013	Mus musculus aczonin (Acz), mRNA	Human L-type calcium channel beta-1 subunit (OACIAL) / Batis	cds Human L-type calcium channel beta-1 subunit (CACNLB1) gene, exon 13B and isoform beta-1B, complete	ods 1802038272F1 NCI CGAP_Brn64 Homo saplens cDNA done IMAGE:4184321 6'	Home saplens calcium channel gamma 5 subunit (CACNG5) gene, expn 4 and compact	1801104462F1 NIH MGC 14 Homo sapiens cDNA clone IMAGE:3347403 o	hz83d11.x1 NCI_CGAP_Lu24 Homo sapiens oDNA clone IMAGE:3214561 3 Silling to 11:323	_		т-	1	1	$\overline{}$	P61843 ORPHAN NUCLEAR RECEPTION 2010 IN A clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 7472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar to SW:DAX1_HUMAN 17472811.x1 NCI_CGAP_Lu24 Home saplens of the clone IMAGE:3278840 3' similar	P61843 ORPHAN NUCLEAR RECEPT ON DAX-1. I.11. P61843 ORPHAN NUCLEAR RECEPT ON DAX-1. I.11. NOBER STA6246 St. SIMILAR to TR.P78336 P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx086R3.X1 NOL. CGAP_OV39 Homo saplens CDNA clone IMAGE: 2746246 St. SIMILAR to TR.P78336 Xx08784 ST. SIMILAR TO TR.P78336 Xx08784 ST. SIMILA	GUANYLATE KINASE ASSOCIATED PROTEIN: i G01589080F1 NIH_MGC_68 Homo saplens dDNA clone IMAGE:3827776 67	601558060F1 NIH_MGC_58 Home sapiens cDNA clone IMAGE:3827775	he34d06x1 NCL CGAP_KId12 Homo saplens cDNA clone IMAGE: 267 3090 5 9111112	Q921N3 MYOSIN-KHOGAP FIXO LENG MINO SEPIENS CDNA QV4.HT0894-290900-399-410 HT0894 Homo sepiens CDNA	
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Buio	Top Hit Acession No.		T	1		A)207616.1	8	DE /911 /3.1	DESENDRO 1	10048478 NT		U86961.1	}	١	0.0E+00 AF142621.1	BE273983.1	0.0E+00 BE503096.1	0.0E+00 BF569905.1	0.0E+00 AA454642.1	0.0E+00 AFZ17.655.1	0.0E+00.0E6201+4.1		0.0E+00 BE673986.1	0.0E+00 BE673986.1	0.0E+00 AW276760.1	0.0E+00 BF031/42.1	0.0E+00 BF031/42.1	+00 AW470846.1	10 1000
	<u>g</u> + 111	_			0.0E+00	0.0E+00			0.01+00	0.05+00	0.0 1	0.0E+00	0.0E+00	0.0円+00	0.0E+00	0.0E+50	0.0日十00	0.0E+00	١	١	١	0.0				١	1	0.0	5
	Expression Signal		1.03	1.56	1.56	1.19	5.23	121	1.13	6.59	1.67	4.05	4.05	2.28	1.03	3.06	0.88			2.38		22/	0.55	0.55	1.14		1.16		9 0.77
	ORF SEQ ID NO:		31344	31404	31405	31416	31430	31433	31444		31448	31447	31448		L		31486					7 31535	31552	31563	31559	31571	Ш		31699
	0	j Z	18431	18485	18485	18493	18509	18514	18523	18524	18525	18526	18526	L	L.	1_	<u> </u>	L	L	18600	18602	18607	18620	<u> </u>	<u> </u>	1			72 18659
		 ö Z	AFAR	200	2694	2689	5717	6722	6731	6732	5733	6734	6734	R752	8788	6767	877A	5773	5778	5811	5813	5818	5831	583		F847	5847	6989	5872

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#91f10.x1 NCI_CGAP_Pr28 Home saplens cDNA clone IMAGE:2248539 3' similar to TR:Q14839 Q14839 zq81d03.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:648005 5' similar to Homo saplens potassium voltage-gated channel, Shal-related subfamily, member 2 (KCND2), mRNA Homo sapiens KIAA0735 gene product, synaptic vesicle protein 2B homolog (KIAA0735), mRNA Homo sapiens KIAA0735 gene product, synaptic vesicle protein 2B homolog (KIAA0735), mRNA 601677736F1 NIH, MGC_21 Homo sapiens cDNA clone IMAGE:3980200 6' 601677735F1 NIH, MGC_21 Homo sapiens cDNA clone IMAGE:3980200 6' 601677735F1 NIH, MGC_21 Homo sapiens cDNA clone IMAGE:3980200 6' 601677735F1 NIH, MGC_21 Homo sapiens cDNA clone IMAGE:3980200 6' Homo sapiens carcinoembryonic antigen-related cell adhesion molecule 8 (CEACAM8), mRNA zc08h06.r1 Soares_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:321755 5' zc08h06_r1 Soares_parathyrold_tumor_NbHPA Home sapiens cDNA clone IMAGE:321755 5 zr40h01.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:665905 5' similar to zr40h01.r1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:665905 5' similar to SW:YY05_HUMAN P42694 HYPOTHETICAL MYELOID CELL LINE PROTEIN 5. SW:YY05_HUMAN P42694 HYPOTHETICAL MYELOID CELL LINE PROTEIN 6. 601158515F1 NIH_MGC_21 Homo saplens cDNA clone IMAGE:3505323 5' 601512630F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3914238 5' 3355565 5' 601236276F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3608490 5' 801109532F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350622 6' Homo saplens familial mental retardation protein 2 (FMR2) gene, exon 14 601114823F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE AU137772 PLACE1 Homo sapiens cDNA clone PLACE1007201 Human G protein-coupled receptor GPR-9-6 gene, complete cds Top Hit Descriptor Human T cell surface glycoprotein CD-6 mRNA, complete cds Human T cell surface glycoprotein CD-6 mRNA, complete cds QV4-HT0894-290900-399-a10 HT0894 Homo sapiens cDNA QV0-HT0368-090200-099-609 HT0368 Homo saplens cDNA Human neurofibromatosis type 1 (NF-1) mRNA, 3' end of cds IR:G854195 G854195 LEUKOCYTE SURFACE PROTEIN. IL3-CT0220-111199-028-E04 CT0220 Homo saplens cDNA Homo saplens xylosyltransferase II (XT2), mRNA Homo sapiens xylosyltransferase II (XT2), mRNA MI-2 PROTEIN.; EST_HUMAN EST_HUMAN HUMAN EST_HUMAN EST_HUMAN **EST_HUMAN** EST HUMAN Top Hit Database HCMAN EST_HUMAN EST HUMAN EST HUMAN EST_HUMAN EST HUMAN EST_HUMAN EST_HUMAN EST HUMAN Source EST 9789986 NT 11426367 NT 11545913 NT **Fop Hit Acession** 11433071 0.0E+00 AW752848.1 BE901608.1 0.0E+00|AA193506.1 0.0E+00 BE889610.1 0.0E+00 AU137772.1 BE257173.1 0.0E+00 AF012618.1 BE280197.1 AA193506.1 BE156561.1 0.0E+00 AA204740.1 0.0E+00|AI686048.1 0.0E+00 BE901608 ġ BE901608. 0.0E+00 W33069.1 0.0E+00 W33069.1 0.0E+00 U45982.1 U34625.1 0.0E+00|M38107.1 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 0.0E+00 (Top) Hit BLAST E Most Simila 3.33 10.66 2.6 16.77 0.85 0.85 2.51 66.0 4.52 3.97 Expression Signal 31600 31605 31609 31639 31642 31665 31690 31839 31604 31620 31641 31644 31664 31840 ORF SEQ 31661 31809 31875 ÖΝΟ 18659 18668 18675 18778 18924 SEQ ID 18664 18664 18690 25081 18710 18710 18730 18788 18845 18873 18872 18907 1891 5872 5878 5882 5905 5908 5908 5909 59.28 59.48 5948 6147 5890 5923 6007 8046 8046 6095 6095 6095 <u>8</u> SEQ ID 5878 5879 5909 6063 5926 5997 9909 5987 6133 ö

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Single Exon Probes Expressed to Brain

					Sir	gle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6151			1.53	9.0E	+00 135930.1	Į,	Human anion exchanger (AF1) gens awar 1.30
6169			1.22	0.0E	+00 BE797385.1	EST HUMAN	1801587971F1 NIH MGC 7 Homo saniens cDNA class NA GE-2042222 51
6159	18936	31904	1.22	0.0E	+00 BE797385.1	EST_HUMAN	801587971F1 NIH MGC 7 Homo sapiens cDNA clone IMAGE:3842328 5
. 6170	18947	31919	0.67	0.0E+00	+00 AI198025.1	EST HUMAN	qi50b11.x1 NCI_CGAP_Brn25 Homo saplens cDNA clone IMAGE:1869901 3' sImilar to TR:Q12838 Q12838 TFIIIC ALPHA SUBUNIT:
6170	18947	31920	0.57	0.0E+00	0.0E+00 Al198025.1	EST HUMAN	qi50b11 x1 NCL CGAP_Brn26 Hamo sapiens cDNA clone IMAGE:1859901 3' similar to TR:Q12838 Q12838 TFIIIC ALPHA SI BI INIT :
6172		31921	0.85	0.0E+00	0.0E+00 BF357123.1	EST HUMAN	MR0-HT0923-220800-102-b05 HT0923 Home and me and me
6180	i	31931	1.08	0.0E+00	11435630 NT	LN	Homo septens pentide transporter 3 // OCE 2001 - DATA
6189	18966	31939	0.65	0.0E+00	D55649	LN	Human mRNA for alpha mannocidese II Inches and the state of the state
6207	18982	31961	1.03	0.0E+00	+00 AW178142.1	EST HUMAN	II.3.HT0082-010999-014-A04-HT0082-Home senions - ANIA
6228		31978	0.66	0.0E+00	+00 BE674544.1	ļ	7e02c12.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:3281302.3' similar to 9W:Y176_HUMAN O14881 HYPOTHETICAL DEDTENNING NAMED CONTROLLED TO SAPERIOR OF SAPERIOR SA
6232		31983	1.33	0.0E+00	7662039 NT		Homo seriens KIAA0285 cene novelust (VIA A02551 - ENIA
6246			8.59	0.0E+00	+00 AV650020.1	T HUMAN	AV650020 GLC Homo saniens CINIA dina CI CCATho 3"
6254		32003	3.13	0.0E+00		Т	UI-HF-BLO-acc-d-12-0. II st NIH MGC 37 Home contact ability of contact at
6257		32008	6.26	0.0E+00	+00 H01255.1	Г	1/27b03.r1 Soares placente Nb2HP Home sentens cDNA clone IMAGE: 41000 #1
6269		32019	1.6	0.0E+00	+00 X15377.1	LN	Human dene for the light and heavy-thens of mail occasions wind introduced to the light and heavy-thens of mail occasions.
6271	19044	32021	0.65	0.0E+00	5.1		8814607.rt Soares NHHMPu S1 Home septens cDNA clone MAGE:817352 F
6272		32022	1.3	0.0E+00	-00 AI612841.1	EST HUMAN	1257408.XI NCI_CGAP_OWS Homo saptems CDNA clone IMAGE:222887 3's imiliar to SW:NTCS_HUMAN P58798 SODII M. AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE TO AND CHI OBIDE DEPENDENT CONTINUE TO AND CHI OBIDE TO AND C
6278		32028	4.71	0.0E+00		i	601305388F1 NIH MGC 39 Homo sanlars child charallity Colorada all
6278		32029	4.71	0.0E+00	0.0E+00 BE735989.1	Т	60130538BF1 NIH MGC 39 Home sapiens cDNA close IMA GE-283984 ET
6282	19055	32035	0.86	0.0E+00	0.0E+00 AW748596.1	Г	MR0-BT0264-221199-002-111 BT0264 Homo sanions - DNIA
6282	. !	32036	0.86	0.0E+00	-00 AW748596.1	Т	MRO-BT0284-221199-002-111 BT0284 Homo capiene child
6283	19056		9.0	0.0E+00[Γ	Т	Homo carllents Achaeles Scribe books of ACC 20
6285		32038	15.59	0.0E+00	Γ	T HUMAN	A 1119245 HEMBA Home sentence and A - LITTLE ASSESSED -
6285		32039	15.59	0.0E+00	0.0E+00 AU119245.1	Ţ	AU19245 HEMBA1 Home sablers CDNA close UEMBA1005500 5
6289	19062	32044	0.8	0.0E+00 E	Γ	Т	601468712F1 NIH MGC 67 Home seniers of NIA Acta MAGE 3974000 5
9230	19063	32045	1.12	0.0E+00	Γ	Г	H.seplens dermine immunodobulin beautable marchia and 122 0
6304	19076	32062	1.52	0.0E+00 AI989483.1		EST HUMAN	WS25c07 XT NCL CGAP GCS Home series china class MA CE success of
6317	19088	32072	6.91	0.0E+00 E	Ĺ	Г	801105344F1 NIH MGC 16 Home septems cDNA clore INVA CE 20077757
6817	19088	32073	6.91	0.0E+00 B		Г	601105344F1 NIH MGC 15 Homo saplens CDNA circa IMA GE-2987893 5
•						1	O CORTIONAL CARRIES CARRIES CONTROL

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					Ę	אם בעחור ויער	Single Extended Expressed in Diam
Probe SEQ ID NO:	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
6355	19125	32119	0.58	0.0E+00	BF057438.1	EST_HUMAN	7k43h05.x1 NCI_CGAP_Ov18 Homo sepiens cDNA clone IMAGE:3478496 3' similar to TK:014553 O14553 R31240_1;
888	1		1.89	0.05+00		EST_HUMAN	UI-HF-BL0-aco-h-02-0-UI.r1 NIH_MGC_37 Homo sapiens cDNA clone IMAGE:3059831 5
6388		32158	1.89		AW406348.1	EST_HUMAN	ULHF-BL0-aco-h-02-0-U.r1 NIH_MGC_37 Homo sapiens cuinA cigne IMAGE: 3003331 3
8418	19186		0.79		AV719444.1	EST_HUMAN	AV718444 GLC Hamo sapiens cDNA clone GLCEHCUS 3
6427	19195		0.98		BE898340.1	EST_HUMAN	601681150F1 NIH MGC 9 Homo sapiens cUNA clone inwace: 3831301 3
6427	19195	32192	0.98	0.0E+00	BE898340.1	EST_HUMAN	601681150F1 NIH MGC 9 Homo sapiens cuiva done liwavet 335150 1
6430	19198	32195	2.24	0.0E+00	AF190860.1	N	Homo septens low voltage-activated 1-type calcium channel apna 16 space varient cavina (Oncond) mRNA, complete cds
8433		1		0.0E+00	11420658 NT	TN	Homo sapiens transformation/transcription domain-associated protein (TRKAP), mKNA
9440	l			0.0E+00	AW163640.1	EST HUMAN	au96h08.y1 Schnelder fetal brain 00004 Homo saplens cDNA clone IMAGE::2784159 5' similar to TR:O15390 O15390 GT24, [3] TR:O43840 TR:O43206;
9440	1	İ				EST_HUMAN	au96h08.yf Schneider fetal brain 00004 Homo sapiens cDNA done IMAGE:2784159 5' similar to TR:O15390 O15390 GT24, [3] TR:O43840 TR:O43206;
6444	1	32208	76.0	0.0E+00	W37163.1	EST_HUMAN	علايات الماسية والماسية الماسية الماسية الماسية الماسية الماسية الماسية الماسية الماسية الماسية الماسية الماسي SW: ZNA5_HUMAN Q02386 ZINC FINGER PROTEIN 45 ;
6444		_			W37163.1	EST_HUMAN	zb20e06.r1 Soeres_fetal_lung_NbHL19W Homo sepiens cDNA clone IMAGE:302626 5' similar to SW:ZN45_HUMAN 002386 ZINC FINGER PROTEIN 45;
6459	I		1.08			EST_HUMAN	601589371F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3943504 5
6466	19233	32233	5.81		_	EST_HUMAN	601587561F1 NIH_MGC_7 Homo sapiens cDNA crone IMAGE:3941047 3
6467	l					EST_HUMAN	QV1-GN0065-140800-318-h02 GN0005 Home sapiens colve-
6467			0.56			EST_HUMAN	QV1-GN00065-140800-310-muz GN0000 name seprens convo
6471	Ш					EST_HUMAN	601512038F1 NIH MGC_/ 1 nomo sapiens cours cione involuces 3311 S
6471						ESI_HUMAN	001012030FT WIT MINO. 7 11 IQUID SQUARES CONTROLLES CON
6480					124493.1	LN.	Human amgen OUZ/ gene, excits 1-2
6485	19262	32251			AL163204.2	<u>₽</u>	Homo sapiens chromosome z1 segment noz roout
6485	1	32252	1.98	L	AL16320	NT	Homo saplens chromosome 21 segment H521C404
649	1	Ì	4.06	0.0E+00	TN 6005983 NT	NT	Homo saplens zona peliucida glycoprotein 3A (sperm receptor) (2F3A), mixivA
	1		22,	_	A 1838442 4	HOT HIMAN	#31f11 x1 NCI_CGAP_GC6 Homo saplens dDNA clone IMAGE:2242413 3 similar to 5W :WN I 3_MOUSE. P17553 WNT-3 PROTO-ONCOGENE PROTEIN PRECURSOR: ;
2484	ı	1				LN LN	Homo sapiens zinc finger homeodomain protein (ATBF1-A) mRNA, complete cds
6495	_1				1.32632.1	EST LINAN	7w52c03 rt Soares total fetus Nb2HF8 9w Homo saplens cDNA clone IMAGE:773688 5
6507	_	32273		1	AA454504. I	NONE FOLLOWING	ANABRA17F1 NIH MGC 57 Homo septems cDNA clone IMAGE:4103693 5
6520		1			JBF21/200.1	TOUNDIA TOU	INV. annova angon 278-n6 Bunda Home septems cDNA
6523	19289	32283	1.82	0.0E+00) BE925875.1	EST HOMAIN	יינים בשלים היינים ביי

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ORF SEQ Expression (Top) Hit Top Hit Acession Database ID NO: Signal BLASTE No. Source	32487 1.08 0.0E+00 11431474 NT	32490 0.6 0.0E+00 BE313075.1 EST_HUMAN	32491 0.6 0.0E+00 BE313075.1 EST_HUMAN	32505 2.69 0.0E+00 BF569905.1 EST_HUMAN	2.32 0.0E+00 J03069.1 NT	32573 3.52 0.0E+00 AF217289.1 NT	32574 3.52 0.0E+00 AF217289.1 NT	32575 1.18 0.0E+00 M38113.1 NT	30571 3.2 0.0E+00 11420775 NT	30572 0.74 0.0E+00/AI419969.1 EST HUMAN	30573 0.74 0.0E+00 AI419969.1 EST HUMAN	30576 0.78 0.0E+00 BE256708.1 EST_HUMAN	30546 0.58 0.0E+00 BE904955.1 EST_HUMAN	30514 1.05 0.0E+00 AU118478.1 EST_HUMAN	30518 8.08 0.0E+00 BE262941.1 EST_HUMAN	30519 2.26 0.0E+00 Z37976.1 NT	30520 2.26 0.0E+00 Z37976.1 NT	30521 3.26 0.0E+00 AF257737.1 NT	30522 3.26 0.0E+00 AF257737.1 NT	30527 1.06 0.0E+00 AF310105.1 NT	32681 1.03 0.0E+00 BE762770.1 [EST_HUMAN	32687 2.37 0.0E+00 BF569905.1 EST_HUMAN	32692 4.53 0.0E+00 L01978.1 NT	32696 0.79 0.0E+00 AW 502362.1 EST_HUMAN	32697 0.79 0.0E+00[AW502362.1 [EST_HUMAN	32704 0.7 0.0E+00 AL039581,1 EST_HUMAN	32705 0.7 0.0E+00 AL039581.1 [EST_HUMAN	32711 5.87 0.0E+00 BF306996.1 EST_HUMAN 601889823F1 NIH_MGC_17 Homo sapiens dDNA clone IMAGE:4123948 5	32715 2.33 0.0E+00 U41302.1 NT	32474 1.18 0.0E+00 AL049784.1 NT	32754
	32487	32490	32491	32505		32573	32574	32575	30571	30672	30573	30576	30546	30514	30518	30519	30520	30521	30522	30527	32681	32687	32692	32696	32697	32704	32705	32711	32715	32474	32754
SEQ ID NO:	19468	19468	ı	1	L	L	L	L.	1	L.		L	L	L	L	L	L	L		L	19637	19642		L	_	<u></u>	L	19665	19669	1	ŀ
Probe SEQ ID NO:	6805	6807	6807	6822	6837	6845	6845	6846	6858	6859	6859	6863	6874	6884	6887	8888	8889	6889	6889	6894	689	6904	8069	6913	6913	6922	6922	6369	6934	6972	700

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	Homo sapiens mRNA for vascular cadherin-2, complete cds	AU137738 PLACE1 Hamo sapiens cDNA clane PLACE1007120 5'	AU137738 PLACE1 Homo sapiens cDNA clone PLACE1007120 5'	EST366876 MAGE resequences, MAGC Homo sapiens cDNA	601113958F1 NIH_MGC_16 Homo saplens cDNA clone IMAGE:3354586 5'	Human type VI sodium channel alpha polypeptide (SCN4A) gene, exon 14	Homo sapiens mRNA for KIAA0466 protein, partial ods	Homo sapiens mRNA for KIAA0488 protein, partial cds	AU133213 NT2RP4 Homo saplens cDNA clone NT2RP4001556 5'	Homo saplens membrane protein CH1 (CH1), mRNA	EST182818 Jurkat T-cells VI Homo sapiens cDNA 5' end	AU143706 Y79AA1 Homo saplens cDNA clone Y79AA1002365 5'	Homo sapiens netrin 1 (NTN1), mRNA	801431819F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3817164 5	601431819F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3917164 6'	Homo sapiens keratin 12 (KRT12) gene, complete c'ds	Homo saplens keratin 12 (KRT12) gene, complete cds	Homo sepiens vitamin D (1,26- dihydroxyvitamin D3) receptor (VDR), mRNA	Homo sapiens vitamin D (1,25- dihydroxyvitamin D3) receptor (VDR), mRNA	Homo sapiens voltage-dependent calcium channel alpha 1G subunit isoform ae (CACNA1G) mRNA, complete cds	qc67e07.x1 Soares_placenta_8tx9weeks_ZNbHP8tx9W Homo sapiens cDNA clone IMAGE:1714644 3' similar to SW:ARSD_HUMAN P51689 ARYLSULFATASE D PRECURSOR ;contains element HGR repetitive element;	qc67a07.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo saplens cDNA clone IMAGE:1714644 3' similar to SW:ARSD_HUMAN P51689 ARYLSULFATASE D PRECURSOR ;contains element HGR	Homo saplens candidate taste receptor T2R9 cene, complete cds	Homo saplens candidate taste receptor T2R9 gene, complete cds	Homo sapiens myosin, heavy polypeptide 8, skeletal muscle, perinatal (MYH8), mRNA	Homo sapiens myosin, heavy polypeptide 8, skeletal muscle, perinatal (MYH8), mRNA	602035089F1 NCI_CGAP_Brn64 Homo sapiens cDNA clone IMAGE:4182839 5'	zn60f09.r1 Stratagene muscle 937209 Homo saplens cDNA clone IMAGE:562601 5' similar to TR:0806562 G806562 NEBULIN.;
gle Exon Prob	Top Hit Database Source	F	EST_HUMAN		HUMAN		F		LN L	EST_HUMAN /		Т	EST_HUMAN		Г	EST_HUMAN					TN 0	EST HUMAN	NAME OF THE PARTY	Т				EST_HUMAN 6	EST_HUMAN G
Sin	Top Hit Acesslan No.	00 AB026893.1	0.0E+00 AU137738.1			0.0E+00 BE254103.1	DO LO1973.1				28081	0.0E+00 AA312125.1	0.0E+00 AU143706.1	883		0.0E+00 BE891286.1	0.0E+00 AF137286.1	0.0E+00 AF137286.1	11436699 NT	11436699 NT	0.0E+00 AF227744.1	0.0E+00 A1128344.1	0 0F+00 A128344 1	T	0.0E+00 AF227135.1	11426392 NT	0.0E+00 11426392 NT		0.0E+00 AA128453.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05	0.0E+00 AF227135.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	0.65	1.07	1.07	1.2	6.0	0.98	0.64	0.64	2.73	0.95	0.56	2.57	0.94	1.32	1.32	2.64	2.54	5.01	5.01	0.65	37.67	37.67	0.68	0.66	4.65	4.65	15.23	2.85
	ORF SEQ ID NO:	32755	32781	32762	32768	32769	32781	32791	32792	32799	32818	32820		32825	32834	32835	30495	30496	32876	32877	32891	32911	32912	32914	32915	32918	32919		32921
	SEQ ID NO:	19700	19705	19705		il			19732	19738	19753	19755	19760	19761	19770	19770	17981	17981	19810	19810	19824	19843	19843	19845	19845	19848	19848	19850	19852
	Probe SEQ ID NO:	7008	7013	7013	7019	7020	7033	7041	7041	7047	7062	7084	7069	7070	7079	7079	7100	7100	7122	7122	7137	7166	7156	7158	7158	7161	7161	7163	7165

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Top Hit Descriptor	DKFZp434B0226_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434B0228 5'	DKFZp434B0226_r1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434B0228 5'	601174578F1 NIH_MGC_17 Hamo saplens cDNA clone IMAGE:3529794 5'	Homo sapiens hypothetical protein (FLJ20281), mRNA	AU118607 HEMBA1 Homo sapiens cDNA clone HEMBA1003969 5'	Homo sapiens ankyrin 1 (ANK1) mRNA, complete cds	Homo sapiens ankyrin 1 (ANK1) mRNA, complete cds	Homo sapiens adlican mRNA, complete cds	H.sapiens DNA for ZNGP2 pseudogene, exon 4	Human P2x1 receptor mRNA, complete cds	Human P2x1 receptor mRNA, complete cds	EST388573 MAGE resequences, MAGD Homo saplens cDNA	7e80h08.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:3223167 3' similar to gb:M54911_ma1 IG HEAVY CHAIN PRECURSOR V-II REGION (HUMAN);	EST382586 MAGE resequences, MAGA Homo saplens cDNA	AF001543 Human cDNA (Chandrasekharappa,S.C.) Homo sapiens cDNA clone kappa_200	AF001543 Human cDNA (Chandrasekharappa,S.C.) Homo saplens cDNA clone kappa_200	AF001543 Human cDNA (Chandrasekharappa,S.C.) Homo sapiens cDNA clone kappa_200	Human BTF3 protein homologue gene, complete cds	601302679F1 NIH_MGC_21 Homo sapiens aDNA alone IMAGE:3637434 5'	UI-HF-BK0-aas-g-07-0-UI.r1 NIH_MGC_38 Hamo sapiens cDNA clone IMAGE:3054924 5'	ym88h10.r1 Soares adult brain N2b4HB55Y Homo sapiens cDNA clone IMAGE:166051 5'	xb39a05.y1 NCI_CGAP_Lu31 Homo sapiens cDNA clone IMAGE:2578640 5' similar to TR:Q08050 Q08050 HNF3/FH TRANSCRIPTION FACTOR GENESIS ;	AU117563 HEMBA1 Homo sapiens cDNA clone HEMBA1001661 5'	Homo sapiens glucagon-like peptide 2 receptor (GLP2R), mRNA	MR0-AN0083-270900-004-t07 AN0083 Homo saplens cDNA	Homo sapiens zinc finger homeodomain protein (ATBF1-A) mRNA, complete cds	601889823F1 NIH_MGC_17 Hamo sapiens cDNA clone IMAGE:4123948 5'	601889823F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4123948 5'	AU118767 HEMBA1 Homo sapiens cDNA clone HEMBA1004314 6'	cn17d05.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn17d05 random
Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	NT	NT	NT	NT	NT	NT	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	AL079497.1	AL079497.1	BE295499.1	11427965 NT	AU118607.1	AF005213.1	AF005213.1	AF245505.1				AW956503.1	BF672445.1		AF001543.1	AF001543.1	AF001543.1	M90354.1	BE408283.1	AW 402542.1	R87430.1	AW239326.1	AU117553.1	11427135 NT	BF229235.1	L32832.1	BF306996.1	BF306996.1	AU118767.1	AI752561.1
Most Similar (Top) Hit BLAST E Value	0.05+00	0.0E+00	0.0E+00						0.0E+00		0.0E+00	0.0E+00	00+30			0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_				0.0E+00	0.0E+00
Expression Signal	7.0	0.7	1.09	1	1.42	1.99	1.99	0.87	8.04	8.51	8.51	0.86	0.56	2.52	0.57	0.57	0.57	0.56	96.0	9.0	1.43	1.88	1.31	3.67	0.58	0.67	1.18	1.18	0.92	4.16
ORF SEQ ID NO:	32927	32928		32970		32973	32974		32989	32891		$\Big _{-}$	33000			33034	33035					33083		33101		33133	33154	33155	33166	33223
	56	19856	19893	19895	19898	19899	19899	19911	19917	19919	19919	19932	1007	19935	19957	19957	19957	19975	19976	19988	20005	20006	20023	20025	20046	20052	20075	20075	20083	20133
SEQ ID NO:	7170 19856	7170 198	7208 19	7210 19		7214 16				ŀ		7247		1		ı	7273 1	١ ١				<u> </u>		1	l	7372	l	7397	7406 2	7460

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TOP HIT Deachase Source EST_HUMAN INT Homo septens dynactin 1 (DCTN1) gene, alternatively spliced products, exons 7 through 32 and complete of the caspiens of process of through 32 and complete of the caspiens of process of through 32 and complete of the caspiens of process of through 32 and complete of thron septens of process of through 32 and complete of thron septens of process of through 32 and complete of thron septens of process of through 32 and complete of thron septens of through 100 through 32 and complete of thron septens of through 100 through 32 and complete of thron septens of through 100 through 32 and complete of thron septens of through 100 through 32 and complete of thron septens of through 100 through 32 and complete of thron septens of through 100 throug	TOP HIT DATABASE SOUTCE SOUTCE SOUTCE STANMAN ES	Top Hit Acess No. No. AF084205.1 AF084205.1 UT4315.1 UT4315.1 UT4315.1 AW 672785.1 AW 672785.1 AR25504.1 AR25504.1 AR25504.1 AR25504.1 AR25504.1 AV069274.1 AV069274.1 BE738670.1 BE738670.1 BE738670.1 BE738670.1		Signal 1.13 1.14 1.17 1.19 1.09 5.87 5.87 6.31 6.31 6.31 1.18	33324 33330 33330 33343 33344 33360 33360 33360 33377 33377 33377 33377 33476 33409 33416 33418 33418	Exen SEQ ID NO: 20133 20239 20239 20276 20276 20276 20276 20276 20276 20276 20276 20276 20276 20370 203710 203710 203710 20370	Probe SEQ ID NO: 7460 T535 T535 T557 T570 T570 T570 T699 T699 T699 T699 T699 T699 T699 T69
AU120424 HEMBB1 Homo saplens cDNA clone HEMBB1000665 6'	EST_HUMAN	AU120424.1	0.0E+00	0.71	33422	20311	7847
TO TAXA LENDED I TO CAPACITO CONTROL TO TAXA LICENTED TO	101	T		12.0	33422		7847
AI HOMASA HEMBER I Dome conjune CDNA alone DEMBERONGEE E!	DOT HIMAN	Γ		0.71	33421	Ш	7647
Homo sapiens atrophin-1 interacting protein 1; activin receptor interacting protein 1 (КІАА0705), mRNA	L L	2461		1.18	33420		7846
Homo saplens atrophin-1 interacting protein 1; activin receptor interacting protein 1 (KIAA0705), mRNA	Ä	6912461	0.0E+00	1.18	33419		7848
601593156F1 NIH_MGC_9 Hamo sapiens cDNA clone IMAGE:3847365 5'	EST_HUMAN		0.0E+00	6.31		- 1	7845
601593156F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3947365 6'	EST_HUMAN		0.0E+00	6.31		- 1	7845
AV758467 BM Homo sapiens cDNA clone BMFBGG05 5'	EST_HUMAN	-	0.0E+00	1.13		-	785 55
Homo saplens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA	LN	1848	0.0E+00	6.26		L	7836
cr42e09.x1 Jia bone marrow stroma Homo saplens cDNA clone HBMSC_cr42e09 31	EST_HUMAN	7	0.0E+00	0.97		$_{-}$	783
cr42e09.x1 Jia bane marrow stroma Homo sapiens cDNA clone HBMSC_cr42e09.3'	EST_HUMAN		0.0E+00			_1	7633
AU128622 NT2RP2 Homo saplens cDNA clone NT2RP2005913 6'	EST_HUMAN	AU129622.1	0.0E+00		1		<u></u>
601885465F1 NIH_MGC_57 Homo sapiens cDNA done IMAGE:4103729 5'	EST HUMAN		0.0E+00	5.87		┚	7604
za86e05.s1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:299456 3'	EST_HUMAN	ĺ	0.0E+00	1.09	ļ		7699
Homo sapiens transient receptor potential channel 5 (TRPC5), mRNA	L	3912735	0.0E+00	1.51		_1	7594
wb17g05.x1 NCL_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2305976 3' similar to TR:076363 075363 AIBC1.;	EST_HUMAN	AI825504.1	0.0E+00	1.97			7586
wb17g05.x1 NCI_CGAP_GC8 Homo saplens cDNA clone IMAGE:2305976 3' similar to TR:075363 075363 AIBC1.;	EST_HUMAN	AI825504.1	0.0E+00	1.97		i	7586
ba01e06.y1 NIH_MGC_7 Homo septens cDNA clone IMAGE:2823108 6' similar to SW P101_PIG 002898 PHOSPHATIDYLINOSITOL 3-KINASE REGULATORY SUBUNIT:	EST_HUMAN	AW672785.1		2.28			7570
ba01e06.y1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:2822108 6' similar to SW:P101_PIG 002698 PHOSPHATIDYLINOSITOL 3-KINASE REGULATORY SUBUNIT;	EST_HUMAN	AW672785.1					7570
Homo sapiens sema domain, seven thrombospondin repeats (type 1 and type 1-like), transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 54 (SEMA5A), mRNA	LΝ						7557
HSU74315 Human chromosome 14 Homo sapiens cDNA clone 1-4	EST_HUMAN	U74315.1					7543
Homo sepiens dynactin 1 (DCTN1) gene, alternatively spliced products, exons 7 through 32 and complete cds	NT	AF064205.1					7535
Homo saplens dynactin 1 (DCTN1) gene, alternatively spiloed products, exons 7 through 32 and complete cds	NT	AF064205.1		,			7535
on17405.x1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clone NHTBC_cn17405 random	EST_HUMAN	AI752561.1					7460
	Top Hit Database Source	Top Hit Acession No.	Most Similar (Top) Hit BLAST E Value	Expression Signal			Probe SEQ ID NO:
ses Expressed in Brain	gle Exon Pro	Sin					

PCT/US01/00667

WO 01/57275

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Probe SEQ ID NO:	Exan SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
7680	20344	33457	1.81	0.0E+00	BE787610.1	EST_HUMAN	601481713F1 NIH_MGC_88 Hamo sapiens cDNA clone IMAGE:3884288 5'
7720	L			0.0E+0	0 W52673.1	EST_HUMAN	zc90f10.rl Pancreatic Islet Homo sapiens cDNA done IMAGE:338443 5'
734	1		0.56	0.0E+0	0 AW 402332.1	EST_HUMAN	UI-HF-BK0-aal-b-08-0-UI.r1 NIH_MGC_36 Homo sapiens cDNA clone IMAGE:3053915 5
7735		33515		0.0	0 AA760692.1	EST HUMAN	nz13e08.s1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1287638 3' similar to gb:U01828 MICROTUBULE-ASSOCIATED PROTEIN 2 (HUMAN);
77.9 E				00-400	0 A A 760692 1	FST HUMAN	nz13a08.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1287638 3' similar to gb:U01828 MICROTUBULE-ASSOCIATED PROTEIN 2 (HUMAN);
7752	1			0.0E+0	0 AU133187.1	EST HUMAN	AU133187 NT2RP4 Homo sapiens cDNA clone NT2RP 4001507 5'
7812	_				BE313013.1	EST_HUMAN	601150347F1 NIH_MGC_19 Homo seplens cDNA clone IMAGE:3503050 5'
7824					0.0E+00 AA149791.1	EST_HUMAN	zo01c06.r1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:566410 5'
7837		33659	0.84	0.0E+00	0.0E+00 BF026628.1	EST_HUMAN	601672310F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3855131 5'
7849	20544		0.45		0.0E+00 AA017021.1	EST_HUMAN	ze33h08.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:360831 5'
7866	ı	33688			BE736046.1	EST HUMAN	601305658F1 NIH_MGC_39 Homo saplens cDNA clone IMAGE:3639903 5
7881			10.48	ı	0.0E+00 M34872.1	NT	Human amyloid-beta protein (APP) gene, exon 11
7881	L	33704			0.0E+00 M34872.1	ΙN	Human amyloid-beta proteth (APP) gene, exon 11
			1		A 100 14 4	MANAGE FOR	bb34d02,1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:2985123 5' similar to TR:064652 064652 E1782 26 PROTEIN :
808/	20004	\$5/55	0.74	1	C.UETON AWO! TOO!.	NUMBER 183	1. COLOR A MILL MCC. 40 Dame continue CDNA close IMAGE-2085433 Ff cimilar to TR-OR4852 OR4852
2809	20604	33735	0.74		0.0E+00 AW674581.1	EST_HUMAN	DD3402.71 NIH_MCC_10 Home septens curva cigle invace_2963123 3 Similar to 11,004322 CO0322 F17K2.26 PROTEIN.;
	l	_			0 0E-00 & \$307551 1	NAMILH FAT	z/81b04.r1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:728719 5' similar to TR:G300482 c300482 POI =REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT) ;
7018	20813			0.05+00	0.0E+00 AW387131.1	EST HUMAN	MR0-ST0031-061099-003-a11 ST0031 Homo saplens cDNA
7821	1				0.0E+00 AB020691.1	본	Homo sapiens mRNA for KIAA0884 protein, partial cds
7922		33744		İ	0.0E+00 AU142402.1	EST_HUMAN	AU142402 Y79AA1 Homo saplens cDNA clone Y79AA1000277 5'
7926	L	L	1.63		0.0E+00 BE388421.1	EST_HUMAN	601285550F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607237 5
7926				l	0.0E+00 BE388421.1	EST_HUMAN	601285550F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607237 5
7942	20637	33764		l	0.0E+00 W95278.1	EST_HUMAN	ze03d01.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:358081 5
7942				l	0.0E+00 W95278.1	EST_HUMAN	ze05d01.r1 Soares_fetal_heart_NbHH19W Homo saplens cDNA clone IMAGE:358081 5
7944	ı		689		BF673096.1	EST HUMAN	602153008F1 NIH_MGC_81 Homo seplens cDNA clone IMAGE:4294128 5
7948	L.	1	0.67	l	AU134114.1	EST_HUMAN	AU134114 OVARC1 Homo sapiens cDNA clone OVARC1001296 5'
7962	L.	7 33782	0.95		BF525534.1	EST_HUMAN	602063632F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4212727 5
7962		33783			0.0E+00 BF525534.1	EST HUMAN	802069632F1 NC CGAP Bm64 Homo saplens cDNA clone IMAGE:4212727 5
7992	20687	L	1.59		AL120124.1	EST_HUMAN	DKFZp761P092_r1 761 (synonym: hamy2) Homo saplens cDNA clone DKFZp761P092 5
7992		33814			AL120124.1	EST_HUMAN	DKFZp761P092_r1 761 (synonym: hamy2) Homo saplens cDNA clone DKFZp761P092 5

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Table 4

Single Exon Probes Expressed in Brain	ORF SEQ Expression (Top) Hit Top Hit Acession Database ID NO: Signal BLASTE No. Source Source	1.32 0.0E+00 BE877893.1 EST HUMAN (601485284F1 NIH MGC 69 Home sanises cDNA clear MADE 38827772 gr	0.0E+00 AW 500549.1 EST_HUMAN	33888 16.05 0.0E+00 AW157233.1 EST HUMAN TR:080463 060463 TYPE-2 PHOSPHATIDIC ACID PHOSPHOHYDROLASE 141.	00 AW072395.1 EST HUMAN	1.09 0.0E+00 11421722 NT	1.07 0.0E+00 W01616.1 EST HUMAN	1.22 0.0E+00 BE745697.1 EST HUMAN	1.22 0.0E+00 BE745597.1 EST HUMAN	0.0E+00 AJ271736.1 NT	33980 0.95 0.0E+00 Al367350.1 EST_HUMAN KIAA0164 PROTEIN ;		33983 1.19 0.0E+00 Ala856711 FST HIMAN SW-COOT HIMAN PROPERTABLY METALL CONTINUE AL	1.07 0.0E+00 BE563650.1 EST HUMAN	1.07 0.0E+00 BE563650.1 EST HUMAN	0.0E+00 11427235 NT	1.63 0.0E+00 11427235 NT	34023 3.2 0.0E+00 AA403192.1 EST_HUMAN TR:G1304132 G1304132 TPRD ;	34024 3.2 0.0E+00 AA403192.1 EST_HUMAN TR:G1304132 G1304132 TPRD :	4.53 0.0E+00 AA398611.1 EST HUMAN PROHIBITIN (HUMAN):	EST HUMAN	1.17 0.0E+00 AW384874.1 EST HUMAN	0.0E+00 AW384874.1 EST HUMAN	1.88 0.0E+00 BE612586.1 EST_HUMAN	1.88 0.0E+00 BE612586.1 EST HUMAN		1.52
			33882	33888	33906	33924	33927	33929	33930	33943	33980	33991	33983	34009	34010	34020	34021	34023	34024					34092	34093	34110	34111
	Exan SEQ ID NO:		20751	20759	20776	20783	1	20798		20809	20848	20859	20861	20874	20874	20883	20883	20885	20885	20925	20934	20935	20935	20954	20954	20969	20969
	Probe SEQ ID NO:	8035	8057	8065	8082	8039	8102	8104	8104	8115	8154	8165	8167	8180	8180	8189	8189	8191	8191	8231	8240	8241	8241	8280	8280	8275	8276

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	Top Hit Descriptor	wm33a11.x1 NCI_CGAP_Ut4 Homo sepiens cDNA clone IMAGE:2437724 3' similer to TR:075457 076457 CT755 CT7	ne25410.s1 NCI_CGAP_Co3 Homo seplens cDNA clone IMAGE:882259 3' similar to TR:G1136434 G1136434 KIAA0187 PROTEIN.;	Homo saplens protocadherin beta 3 (PCDHB3), mRNA	ta04f11.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:2043117 3'	601431238F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3916569 5'	2822701.5prime NIH_MGC_7 Homo sapiens cDNA clone IMAGE:2822701 5	2822701.5prtme NIH_MGC_7 Homo saplens cDNA clone IMAGE:2822701 5	Homo sapiens mitogen-ectivated protein kinase kinase kinase 13 (MAP3K13), mRNA	Homo saplens mitogen-ectivated protein kinase kinase kinase 13 (MAP3K13), mRNA	Human zinc finger protein (ZNF165), gene, exons 2 and 3	Human zinc finger protein (ZNF165), gene, exons 2 and 3	Homo sapiens NESP55, GNAS1 antisense (partial) and XLalphas (partial) genes	H.sapiens mRNA for gamma-glutamytransferase	H.saplens mRNA for gamma-glutamyltransferase	H.saplens mRNA for gamma-glutamyltransferase	Human Immunoglobulin-like transcript-3 mRNA, complete cds	Homo sapiens cep250 centrosome associated protein mRNA, complete cds	Homo sapiens cep250 centrosame associated protein mRNA, complete cds	AU131671 NT2RP3 Homo saplens cDNA clone NT2RP3003016 51	Homo saplens immunoglobulin superfamily, member 2 (IGSF2), mRNA	x46e01.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2707032 3' sImilar to gb:M14123_cds4 RETROVIRUS-RELATED POL POLYPROTEIN (HUMAN);	HUM084C02B Clontech human fetal brain polyA+ mRNA (#6535) Homo saplens cDNA clone GEN-084C02	5.	601236488F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3608709 5'	zt32e04.r1 Soares ovary tumor NbHOT Homo sapiens cDNA clone IMAGE:724062 51	601900571F1 NIH_MGC_19 Homo sapiens cDNA done IMAGE:4129744 5'	Homo saptens keukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 3 // III DB3\	III.H.Ritdr12.0.1 st NCI CGAP Sub3 Home sepiens cDNA clone IMAGE:2717687 3	OLYMPIA COLOR OLYMPIA COLOR CO	
יייי פול	Top Hit Detabase Source	EST_HUMAN	EST_HUMAN	F	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ΝΤ	NT	NT	TN	TN	NT	TN	TN	NT	TN	ΝT	EST_HUMAN	ĻΝ	EST HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	Ŀ	TOT LIMAN	ESI DOMAIN	HOW HE HAVE
Ole III	Top Hit Acession No.	0.0E+00 AI884477.1		6229	0.0E+00 AI580780.1			0.0E+00 AW245765.1	4758695 NT	4758695 NT			1.1					0.0E+00 AF022655.1		AU131671.1	0.0E+00 11426572 NT	00 AW513513.1		D52650.1	0.0E+00 BE378495.1	0.0E+00 AA410545.1	00 BF313946.1	20070777	114420070	+UU AW 1396/3.1	AW1396731
	Most Similar (Top) Hit BLAST: E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U88084.1	0.0E+00	0.0E+00 X98922.1	0.0E+00 X98922.1	0.0E+00 X98922.1	0.0E+00 U82979.1	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	L	0.00	0.0=+00	100+100
	Expression Signal	1.3	1.27	0.59	1.02	1.84	0.72	0.72	2.24	2.24	0.59	0.59	99.0	2.63	2.63	2.63	99.0	0.88	0.88	2.28	0.65	1.92		14.55	4.04	2.58	2.91				1 46
	ORF SEQ ID NO:	34120			34137		34163	34164	34165	34166	34169					ļ	34255			L				34323		١	İ		1	34374	
	SEQ ID	20980	1	20992	20999	21002	21027	21027	21028	21028	l	1	L		L	L		1	1		1	21179		21181	i i	21218	i .	İ	-1.	- 1	21232
	Probe SEQ ID NO:	8286	8283	8238	8305	8308	8334	8334	8335	8335	8339	8339	8404	8409	8409	8409	8424	8465	8465	8468	8483	8487		8480	8520	8528	8528		8535	8540	8540

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					Sin	gle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
8545	21237		0.49		0.0E+00 AI640190.1	EST HUMAN	we30b10x1 NCL_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2299579 3' sImilar to TR:015044 O15044 KIAA0335.
8564			0.76		0.0E+00 BF377897.1	EST HUMAN	CM1-TN0141-250900-439-b08 TN0141 Homo sapiens cDNA
8574	21266	34406		L		TN	Homo sapiens chromosome 21 seament HS21C101
8580			5.89			EST HUMAN	601150051F1 NIH MGC 19 Homo saplens cDNA clone IMAGE 3502838 57
8585	21277	34414	2.51			EST HUMAN	602127664F1 NIH MGC 56 Homo sapiens cDNA clone IMAGE:4284542 51
8585		34415	2.51	0.0E+00		EST_HUMAN	602127684F1 NIH MGC 56 Homo saplens cDNA clone IMAGE:4284542 5
8685	- 1	34416	2.51	0.0E+00		EST_HUMAN	802127664F1 NIH MGC 36 Homo saptens cDNA clone IMAGE:4284542 51
860		34434	0.53	0.0E+00		EST_HUMAN	tk13h11.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2150949 3'
8626	21318	34460	0.85	0.0E+00		EST_HUMAN	AL449770 Homo sapiens fetal brain (Stavrides GS) Homo sapiens cDNA
8631	21323	34464	7 75	004400		HOT LINAAN	or80g02.s1 NCI_CGAP_Lu5 Home sapiens cDNA clone IMAGE:1602194 3' similar to gb:M36072 60S
8637		34472	3.09	0.0E+00	0.0E+00 10947037 NT	LA LA	Home sanlans ankerin 1 andreweits (ANK4) transmit trained at the Bhia
8637		34473	3.09	0.0E+00	10947037 NT	LZ	Homo saplens ankyrin 1, crysticocytic (ANKA) transcript variant 1, mruk Homo saplens ankyrin 1, endhrocytic (ANKA) transcript variant 1, mbily
8860	21362	34499	1.3	0.0E+00	Γ	LN	Home sablens ITGB4 cana for intervity before 4 subjust evens 2.44
8662	21354	34501	1.62	0.0E+00	Γ	EST HUMAN	601156330F1 NIH MGC 21 Homo saciens cDNA clone IMAGE 3130734 F
8672	21364		1.91	0.0E+00	0.0E+00 AV718377.1	EST_HUMAN	AV718377 FHTB Homo sapiens cDNA clone FHTBAAF11 5'
8679	21371	34516	3.33	0.0E+00	0.0E+00 AW337277.1	EST HUMAN	xw73c07x1 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE::2833644 3' similar to gb:X63587
9886		34521	1.12	0.0E+00	Γ	EST HUMAN	AU124051 NT2RM2 Homo sapisns cDNA clone NT2RM2001875 51
8761	21453	34601	1.05	0.0E+00	0.0E+00 AU140704.1	EST HUMAN	AU140704 PLACE4 Homo saplens cDNA clone PLACE4000089 5/
8771	21463	34610	0.86	0.0E+00	3.1	TN	Homo sapiens mRNA for KIAA0454 protein, partial cds
8776	. 1	34614	0.54	0.0E+00 R17132.1		EST_HUMAN	yg09e09.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:31674 5'
8778	21468	34615	0.54	0.0E+00		EST_HUMAN	19909609.r1 Soares Infant brain 1NIB Homo saplens cDNA clone IMAGE:31674 51
8780	21472	34617	4.43	0.0E+00		EST_HUMAN	hf48a09.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE: 2835096 3'
828	- 1	34618	4.43	0.0E+00	AW 592233.1	EST_HUMAN	Hf48a09.x1 Soares_NFL_T_GBC_S1 Home sapiens cDNA clone IMAGE:2835096 31
8816	ı	34852	0.47	0.0E+00	AU128804.1	EST_HUMAN	AU128804 NT2RP2 Homo sapiens cDNA clone NT2RP2004248 5'
8827	ı	34664	1.04	0.0E+00	AV714764.1	EST_HUMAN	AV714764 DCB Homo sapiens cDNA clone DCBAUA08 5'
843 643	21635	34679	2.79	0.0E+00	AL040428.1		DKFZp434C1814_s1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C1814 3'
8843	21635	34680	2.79	0.0E+00/	AL040428.1	EST HUMAN	DKFZp434C1814_s1 434 (synonym: htes3) Homo sapiens cDNA clone DKFZp434C1814 3'
8849	21640	34686	1.17	0.01100	AF133901 1	LΝ	Homo sapiens killer inhibitory receptor 2-2-1 (KIR221) and killer inhibitory receptor 2-2-2 (KIR222) genes, partial are
8851	١.	34689	2.03	0.0E+00,			Homo sapiens mRNA for KIAA1512 problem partial cds
8868	ı	34696	0.65	0.0E+00		T HUMAN	602138483F1 NIH MGC 83 Home septens cDNA clone IMAGE 4274708 F
					1	1	

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					5		יייייייייייייייייייייייייייייייייייייי
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEO ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession 'No.	Top Hit Database Source	Top Hit Descriptor
8860	21551		0.8		0.0E+00 BF058289.1	EST_HUMAN	7K2Bb03.X1 NCI_CGAP_Ov18 Homo sepiens cDNA clone IMAGE:3476692 3' similar to TR:036448 036448 S GAG.;
6888	21580	34720	3.97	Ī	11422857 NT	LΝ	Homo saplens tumor protein p73 (TP73), mRNA
8838	21589	34729	1.19		0.0E+00 K01241.1	TN	Human ig rearranged H-chain epsilon-3 pseudogene, constant region
8905		34737	4.27	0.05+00	0.0E+00 AB020630.1	ΝΤ	Homo sapiens mRNA for KIAA0823 protein, partial cds
8905	L	34738	4.27	0.0E+00	0.0E+00 AB020630.1	TN	Homo saplens mRNA for KIAA0823 protein, partial cds
8940	L	34744	1.79		0.0E+00 AV660739.1	EST_HUMAN	AV660739 GLC Hamo saplens cDNA clone GLCGKG123'
8916		34750	2.88	H	7706638 NT	١. ١	Homo sapiens polycystin-L (PKDL), mRNA
8921		34755			0.0E+00 BE793326.1	EST_HUMAN	601598304F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3942553 5
8922	L					NT	Homo sapiens mRNA for KIAA1251 protein, partial cds
8922		34757		l	0.0E+00 AB033077.1	NT	Homo sapiens mRNA for KIAA1251 protein, partial cds
8934					H73937.1	EST_HUMAN	yu03h08.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:232767 5'
8944	L	34779		0.0E+C	0.0E+00 BE315402.1	EST HUMAN	601141119F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3140740 5
8944	١.	34780	4.57		0.0E+00 BE315402.1	EST_HUMAN	601141119F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3140740 5'
8954		34795	0.48		0.0E+00 BE612721.1	EST_HUMAN	601452582F1 NIH_MGC_66 Home saplens cDNA clone IMAGE:3856100 5'
8954		34796			0.0E+00 BE612721.1	EST_HUMAN	601452582F1 NIH_MGC_66 Homo sapiens cDNA clone IMAGE:3856100 5'
8957	I_		0.45		M89986.1	TN	Human polymorphic loci in Xq28
8929	L	34800	3.68		0.0E+00 X14768.1	FN	Human mRNA for GABA-A receptor, alpha 1 subunit
8980	L_	34820	2.03	0.0E+C	JO A1061395.1	EST_HUMAN	an29e04.x1 Gessler Wilms tumor Homo capiens cDNA clone IMAGE:1700094.3'
	L						wq34q12.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2473150 3' similar to SW:MGB3_HUMAN
8982					0.0E+00 AI954607.1	EST_HUMAN	015480 MELANOMA-ASSOCIATED ANTIGEN B3;
8983					9256595 NT	NT	Homo sapiens protocadherin apha 8 (PCDHA8), mRNA
8000		34840	1.42	0.0E+C	30 AW958311.1	11.1 EST_HUMAN	EST370381 MAGE resequences, MAGE Homo saplens cDNA
9011		34851		0.0E+C	9635487	NT	Human endogenous retrovirus, complete genome
9020				0.0E+(30 AU142662.1	EST_HUMAN	AU142662 Y79AA1 Homo sapiens cDNA clone Y79AA1000678 5
9042	ட				11436995 NT	LN	Homo sapiens MAP-kinase activating death domain (MADD), mRNA
9043	<u> </u>			0.0E+C	JO BE410768.1	EST_HUMAN	601301676F1 NIH_MGC_21 Hamo sepiens cDNA clone IMAGE:3636163 5'
	ĺ			١.			7g97h12.x1 NCI_CGAP_Co16 Home sapiens cDNA clone IMAGE:3314471 3' similar to TR:Q9UH62
9050			7		0.0E+00 BF002024.1	EST_HUMAN	Q9UH62 HYPOTHETICAL 42.5 KD PROTEIN.;
9070	21759		1.1	0.0E+(١	NT	Homo sapiens mRNA for KIAA0578 protein, partial cds
9071		34921	7.72	0.0E+(EST_HUMAN	801589294F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3943463 5
9075	1			0.0E+(EST_HUMAN	RC3-PT0151-290600-011-c05 PT0151 Homo saplens cDNA
9075	ĺ				0.0E+00 BE810292.1	EST_HUMAN	RC3-PT0151-290600-011-c05 PT0151 Homo sapiens cDNA
9078	21767		2.93	0.0E+(30 AU136229.1	EST_HUMAN	AU136229 PLACE1 Homo sapiens cDNA clone PLACE1003804 5

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Single Exon Probes Expressed in Brain	Top Hit Descriptor	601510247F1 NIH MGC 71 Homo saplens cDNA clone IMAGE:3911986 5:	601510247F1 NIH MGC_71 Homo saplens cDNA clone IMAGE:3911988 5'	Homo saplens mRNA for KIAA0594 protein, partial cds	EST50505 Gall bladder I Homo sapiens cDNA 5' end	EST50505 Gall bladder I Homo saplens cDNA 5' end	ba54d08,38 NIH MGC_10 Home saplens cDNA clone IMAGE:2900367 5' similar to TR:060275 060275 KIAA0522 PROTEIN;	ba54d08.y3 NIH_MGC_10 Home sapiens cDNA done IMAGE:2900387 5' similar to TR:080275 080276 KIAA0522 PROTEIN;	ba09f05,y1 NiH_MGC_7 Homo sapiens cDNA clone IMAGE:2823873 5' similar to gb:L35049 Mus musculus Bcl-xL mRNA, complete cds (MOUSE);	be09065.y1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:2823873 6' similar to gb:L35049 Mus musculus Bolad mRNA complete ade (MOI ISE)	802023450F1 NCI CIGAD BINGT Home contens of NA claim MAACE: 44 50200 F1	QV2-HT0698-250700-282-b08 HT0698 Homo saplens cDNA	601455118F1 NIH MGC 66 Hamo sepiens cDNA clane IMAGE:3859035 51	601455118F1 NIH MGC_86 Homo saplens cDNA clone IMAGE:3859036 5'	RC-BT108-040359-032 BT108 Hamo saplens cDNA	Homo saplens leukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 6 (LILRB5), mRNA	Homo sapiens leukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 6 (LILRB5), mRNA	DKFZp434L0120_r1 434 (synonym: htes3) Hamo sapiens cDNA clone DKFZp434L0120 6'	ow60h01.xf Soares_NSF_F8_BW_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:1651249.3' similar to TR:Q14677 Q14677 KIAA0171 PROTEIN.:	801892245F1 NIH_MGC_17 Homo saplens cDNA clone IMAGE:4138068 5	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mRNA	Homo sapiens hypothetical C2H2 zlnc finger protein FLJ22504 (FLJ22504), mRNA	qm09a06.x1 NCI_CGAP_Lu5 Hamo sapiens cDNA clone IMAGE:1881288 3' similar to SW:RL2B_HUMAN P28318 60S RIBOSOMAL PROTEIN L23A. ;	gm09a08.x1 NCI_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1881298 3' similar to SW:RL2B_HUMAN P23316 ets RIBOSOMAL PECTEIN 1 24 A	EST386028 MAGE resequences, MAGC Homo saplens cDNA	Homo saplens polycystic kidney disease 2-like protein (PKD2L) gene, exon 8
le Exon Probes	Top Hit Database Source	EST HUMAN 60	T		EST_HUMAN ES	Г	Bat EST_HUMAN KIA	Bat EST_HUMAN KIA			Т	Т	EST HUMAN 601	Т				EST_HUMAN DKI	EST HUMAN TR:				EST_HUMAN P29	qm0	Т	
Sing	Top Hit Acession No.	0.0E+00 BE883843.1 E				0.0E+00 AA344601.1 E		00 AW673469.1 E	00 BE207063.1 E	0.0E+00 BE207083.1		L			0.0E+00 AI908351.1 E	5803069 NT	5803069 NT	0.0E+00 AL042278.1 E		0.0E+00 BF309962.1 E	0151	11560151 NT			Γ	0.0E+00 AF153466.1 NT
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 /	0.0E+00	0.0E+00 E	0.0E+00	0.0E+00	0.0E+00 AI290909.1	0.0E+00.A	0.0E+00 A	0.0E+00
-	Expression Signal	1.27	1.27	0.62	1.4	1.4	1.13	1.13	1.62	1.62	1.61	2.77	0.88	0.88	0.5	0.81	0.81	1.5	1.28	0.72	2.51	2.51	9.88	88.6	1.99	3.07
	ORF SEQ ID NO:	34935	34936	34953	34957	34958	34998	34999	35031	35032	35260	35093	35201	35202	35209	35211	35212	35137	36171	33524	33527	33528	33531	33532	33533	35117
	Exon SEQ ID NO:	21772				1	21834	21834	21867	21867	22088	21823	22031	22031	22037	22040	22040	21963	21998	20409	20411	20411	20413	20413	20414	21945
	Probe SEQ ID NO:	9083	9083	9102	9108	9108	9164	9164	9188	9198	9209	9244	9277	9277	9283	9286	9286	9596	9331	9338	8340	9340	9342	9342	8343	9370

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
9373	Ι.	35121	99'0	0.05+00	BE885128.1	EST_HUMAN	601510882F1 NIH_MGC_71 Homo sapiens cDNA clone IMACE:3912165 5'
5755	21948	35122	99'0	0.0E+00	BE885128.1	EST_HUMAN	601510882F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912165 5
9382	١.		7.32	0.0E+00	BE255829.1	EST HUMAN	601109942F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350722 5'
9385	1	35219	1.09	0.0E+00	BE781382.1	EST_HUMAN	601466828F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870007 5
9385	22047	35220	1.09	00+30'0	BE781382.1	EST_HUMAN	601466828F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870007 5'
9387	22049	35221	12.62		AW163779.1	EST HUMAN	au86c04.y1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783142 5' sImilar to gb:M36072, 60S RIBOSOMAL PROTEIN L7A (HUMAN);
9409	1		2.98	0.0E+00	BE263191.1	EST_HUMAN	601145054F2 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3160477 5'
9427	ł	35278	4.29	0.0E+00	C06158.1	EST_HUMAN	C06158 Human pancreatic islet Homo saplens cDNA clone hbc5605
9427	l		4.29	0.0E+00	C06158.1	EST_HUMAN	C06158 Human pancreatic islet Homo sapiens cDNA clone hbc5005
9429	L	35282	2.63	0.0E+00	BE746215.1	EST HUMAN	601578683F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3927548 5'
9439	L			L	11437282 NT	NT	Homo saplens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA
9439				L	11437282 NT	NT	Homo saplens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA
9439			2.14		11437282 NT	LΝ	Homo sapiens solute carrier family 21 (organic anion transporter), member 9 (SLC21A9), mRNA
9459			1.44	0.0E+00	BE900549.1	EST_HUMAN	601673425F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3956238 5'
9475	22128		1.01	0.0E+00	AV701829.1	EST_HUMAN	AV701829 ADB Homo sapiens cDNA clone ADBBYH01 5'
9489		35321	2.62	0.0E+00	AF019084.1	NT	Homo sapiens keratin 2e (KRT2E) gene, complete cds
9489				0.0E+00	AF019084.1	NT	Homo sapiens keratin 2e (KRT2E) gene, complete ods
9522		35359	0.94	0.0E+00	BE082977.1	EST_HUMAN	RC2-BT0642-130300-017-g01 BT0642 Homo sapiens cDNA
9541	22194	35379	1.74	0.0E+00	AW 500293.1	EST_HUMAN	UI-HF-BN0-akg-b-12-0-UI.r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3076943 5'
9541	22194	35380	1.74	0.0E+00	AW 500293.1	EST_HUMAN	UI-HF-BN0-akg-b-12-0-UI:r1 NIH_MGC_50 Homo sapiens cDNA clone IMAGE:3076943 51
9550	22203	35386	1,45	0.0E+00	AF029308.1	LN.	Homo sapiens chromosome 9 duplication of the T cell receptor beta locus and trypsingen gene families
9550	22203	35387	1.45		AF028308.1	LN.	Homo saplens chromosome 9 duplication of the T cell receptor beta locus and trypsinogen gene families
9552	ı	35388	69'0	0.0E+00	BE783272.1	EST_HUMAN	601470824F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3874037 5'
9552	22205		69.0	0.0E+00	BE783272.1	EST_HUMAN	601470824F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3874037 5'
9561	_	35400	0.54	0.0E+00	W 56629.1	EST_HUMAN	2d16e11.r1 Soares_fetal_heart_NbHH19W Homo sepiens cDNA done IMAGE:340844 5
9561		35401	0.54	0.0E+0	W 56629.1	EST_HUMAN	zd16e11.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:340844 51
9572	L	35410	1.83	0.0E+00	AB035356.1	NT	Homo sapiens mRNA for neurexin I-apha protein, complete cds
9256	1		8.0	0.0E+00	Ai124780.1	EST_HUMAN	am56a11.x1 Johnston frontal cortex Homo saplens cDNA clone IMAGE:1539548 3'
9578	1			0.0E+00	AW 500526.1	EST_HUMAN	UI-HF-BN0-akj-c-07-0-UI.r1 NIH_MGC_50 Homo saplens cDNA clone IMAGE:3077364 5
9624	22277	35466	1.53	0.0E+0	0 AF009668.1	NT	Multiple scierosis associated retrovirus polyprotein (pol) mRNA, partial cds

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PCT/US01/00667

WO 01/57275

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Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top HIt Database Source	Top Hit Descriptor
9918	22567	35763	2.98		0.0E+00 AF072408.1	NT	Homo saplens killer cell inhibitory receptor KIRCI gene, exons 2, 3, and 4
1266	22569	35765	2.75	0.0E+00	11421001 NT	NT	Homo saplens HEF like Protein (HEFL), mRNA
9921	1	35766		0.0E+00	11421001 NT	TN	Homo sapiens HEF like Protein (HEFL), mRNA
9956	ì			0.0E+00	0.0E+00 AU136637.1	EST_HUMAN	AU136837 PLACE1 Hamo sapiens cDNA clone PLACE1004737 5'
9926		35810	8	0.0E+00	0.0E+00 AU136637.1	EST_HUMAN	AU136637 PLACE1 Hamo saplens cDNA clone PLACE1004737 5
9972	l	L	2.08		0.0E+00 AJ295844.1	NT	Homo sapiens partial RANBP7 gene for RanBP7/Importin7 and partial ZNF143 gene
9972		35826	2.08		0.0E+00 AJ295844.1	NT	Homo sapiens partial RANBP7 gene for RanBP7/Importin7 and partial ZNF143 gene
7766			1.04		0.0E+00 AV695712.1	EST_HUMAN	AV695712 GKC Homo sapiens cDNA clone GKCDXA07 5
2266	22625	35833			0.0E+00 AV695712.1	EST_HUMAN	AV695712 GKC Homo sapiens cDNA clone GKCDXA07 5'
8883	1	35840	0.74		AF072408.1	NT	Homo sapiens killer cell inhibitory receptor KIRCI gene, exons 2, 3, and 4
8882	l		3.11		0.0E+00 AA196387.1	EST_HUMAN	本97h11.r1 Strategene muscle 837209 Homo saplens cDNA clone IMAGE:628197 5
1001	_		-	0.0E+00	0.0E+00 AA131248.1	EST_HUMAN	231f01.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:503545 5
1001	22659			0.0E+00	0.0E+00 AA131248.1	EST_HUMAN	z31f01.r1 Soares_pregnant_uterus_NbHPU Homo septens cDNA clone IMAGE:503545 6
10056					AF179308.1	NT	Homo sapiens KiF4 (KiF4) mRNA, complete cds
10101	1_	35964	0.92		0.0E+00 BE880658.1	EST_HUMAN	601491565F1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3893657 5'
10112	L				0.0E+00 BE730772.1	EST_HUMAN	601570712F1 NIH_MGC_21 Hamo saplens cDNA clone IMAGE:3845403 5'
10112					BE730772.1	EST_HUMAN	601570712F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3845403 6'
10117	<u> </u>		76.0		0.0E+00 AU127403.1	EST_HUMAN	AU127403 NT2RP2 Hamo saplens cDNA clone NT2RP2001212 5
10127	22775				0.0E+00 BE958511.1	EST_HUMAN	601645134F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:3830177 5
10127	上	35989			0.0E+00 BE958511.1	EST_HUMAN	601645134F1 NIH_MGC_56 Hamo sapiens cDNA clone IMAGE:3930177 5'
10144	1_				0.0E+00 BE897487.1	EST_HUMAN	601432317F1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3917453 5'
10154	ı				AA31162	EST_HUMAN	EST182353 Jurkat T-cells VI Homo sapiens cDNA 5' end
10155	22803				4758827 NT		Homo sapiens neurexin III (NRXN3) mRNA
10166					0.0E+00 BE891113.1	EST_HUMAN	601432228F1 NIH_MGC_72 Hamo sapiens cDNA clone IMAGE:3917598 5
10169	22817	36035	1.13		11560151 NT	ΝΤ	Homo sapiens hypothetical C2H2 zinc finger protein FLJ22504 (FLJ22504), mKNA
10179		36041	1.72		AB029290.1	NT	Homo sapiens mRNA for actin binding protein ABP620, complete cds
10180		L		<u> </u>	0.0E+00 BE304522.1	EST_HUMAN	601105459F1 NIH_MGC_15 Homo sapiens cDNA done IMAGE:2987918 5
10180					0.0E+00 BE304522.1	EST_HUMAN	601105459F1 NIH_MGC_15 Hamo sapiens cDNA clone IMAGE:2987918 5'
10187					0.0E+00 AB006590.1	NT	Homo sapiens mRNA for estrogen receptor beta, complete ods
10187	22835		6.02		0.0E+00 AB006590.1	NT	Homo saplens mRNA for estrogen receptor beta, complete cds
					, , , , , ,		219b06.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sepiens cDNA clone IMAGE:450707 3' similar to
10183	_ [_	0.0E+00/AA/0445/.1	ES L'ACIMEN	Himen hate 1 Analorized-transferase mRNA complete cits
10195	١				0.0E+00 MZZZZ1.1	IN I	Human India II-Yakaran ya anininina china kata ilaya kata kata kata kata kata kata kata k
10197	22845	38060	5.45		0.0E+00 BF340331.1	ESI HOMAN	י פטרבטן דיבייטרוייון טוואס סוואין ספואין דיטווען באטרעט פאטרעט באטרעט פאטרעט באטרעט באטרעט באטרעט באטרעט באטרעט

Page 526 of 536 Table 4

Probe Exerce of 1000 Fig. 1000 Proper SEC Disposal of Trop Ht Aceasian Auguste Top Ht Descriptor Top Ht Descriptor SEC ID SEC ID DIA (1972) BLO. SEC ID SEC ID DIA (1972) Top Ht Aceasian Auguste Database Top Ht Descriptor 10197 22820 SSEG ID SEC ID S							פום בייטיים פום	Tobas Explassed II Diain
22845 36061 5.45 0.0E+00 BF34033·1 EST_HUMAN 22870 36082 0.33 0.0E+00 BE897149·1 EST_HUMAN 22800 36110 0.55 0.0E+00 AV716271·1 EST_HUMAN 22800 36110 0.55 0.0E+00 AV716271·1 EST_HUMAN 22800 36143 2.36 0.0E+00 AV716271·1 EST_HUMAN 22830 36149 0.55 0.0E+00 AV716271·1 EST_HUMAN 22830 36149 0.0E+00 AV716271·1 EST_HUMAN 22896 36149 0.0E+00 AV631818·1 EST_HUMAN 22807 36190 0.46 0.0E+00 AV631818·1 EST_HUMAN 22807 3624 2.5 0.0E+00 AV64766.1 EST_HUMAN 22808 3624 2.5 0.0E+00 BE436218·1 EST_HUMAN 22808 36276 2.75 0.0E+00 BE436218·1 EST_HUMAN 23025 36276 0.75 0	Probe SEQ ID NO:		ORF SEQ ID NO:	Expression Signal		Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
22870 36082 0.93 0.0E+00 BE897149.1 EST_HUMAN 22870 36083 0.83 0.0E+00 BE897149.1 EST_HUMAN 22800 36111 0.55 0.0E+00 AV716271.1 EST_HUMAN 22830 36143 2.36 0.0E+00 AV716271.1 EST_HUMAN 22830 36144 2.36 0.0E+00 AV716271.1 EST_HUMAN 22830 36149 0.49 0.0E+00 AV716271.1 EST_HUMAN 22846 36190 0.0E+00 AV716271.1 EST_HUMAN 22896 36190 0.0E+00 AV6172429.1 EST_HUMAN 228970 36234 2.5 0.0E+00 BE436218.1 EST_HUMAN 23020 36234 2.5 0.0E+00 BF436218.1 EST_HUMAN 23021 36236 0.0E+00 BF436218.1 EST_HUMAN 23021 36236 0.0E+00 BF64213.1 EST_HUMAN 23021 36236 0.0E+00 BF64213.1 EST_HUMAN<	10197				0.0E+00	BF340331.1	EST_HUMAN	602037045F1 NCI_CGAP_Brn64 Homo saplens cDNA clone IMAGE:4184939 6'
22870 36083 0.89 0.0E+00 BE897149.1 EST_HUMAN 22800 38110 0.55 0.0E+00 AV716271.1 EST_HUMAN 22800 36141 0.55 0.0E+00 AV716271.1 EST_HUMAN 22830 36148 2.36 0.0E+00 AI631818.1 EST_HUMAN 22836 36149 0.49 0.0E+00 AI631818.1 EST_HUMAN 22846 36190 0.64 0.0E+00 AI631818.1 EST_HUMAN 22846 36190 0.64 0.0E+00 AI631818.1 EST_HUMAN 22846 36190 0.64 0.0E+00 AI631818.1 EST_HUMAN 22846 36190 0.64 0.0E+00 AI631818.1 EST_HUMAN 22847 36190 0.64 0.0E+00 BI6436218.1 EST_HUMAN 2286 36214 2.5 0.0E+00 BI6436218.1 EST_HUMAN 23021 36236 3.25 0.0E+00 BI6436218.1 EST_HUMAN 23081	10222		36082		0.0E+00	BE897149.1	EST HUMAN	801439713F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3924678 6'
22900 36110 0.55 0.0E+00 AV716271.1 EST_HUMAN 22830 36141 0.55 0.0E+00 AV716271.1 EST_HUMAN 22830 36144 2.36 0.0E+00 AI631818.1 EST_HUMAN 22945 36149 0.49 0.0E+00 AI631818.1 EST_HUMAN 22945 36190 0.64 0.0E+00 AI631818.1 EST_HUMAN 22946 36190 0.64 0.0E+00 AI632229.1 EST_HUMAN 22970 36196 0.46 0.0E+00 AU122429.1 EST_HUMAN 22971 36214 2.5 0.0E+00 BF436218.1 EST_HUMAN 22905 36214 2.5 0.0E+00 BF436218.1 EST_HUMAN 23020 36229 2.75 0.0E+00 BF436218.1 EST_HUMAN 23020 36229 2.75 0.0E+00 BE649213.1 EST_HUMAN 23020 36229 2.75 0.0E+00 BE6492120.1 EST_HUMAN 23020 <	10222		36083	0.93	0.0E+00	BE897149.1	EST_HUMAN	801439713F1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3924578 6'
22900 36111 0.55 0.0E+00 AV716271.1 EST_HUMAN 22830 36143 2.36 0.0E+00 AI631818.1 EST_HUMAN 22836 36149 0.26 0.0E+00 AI631818.1 EST_HUMAN 22945 36169 1.52 0.0E+00 AI631818.1 EST_HUMAN 22976 36190 0.64 0.0E+00 AU122428.1 EST_HUMAN 22976 36196 0.46 0.0E+00 AU122428.1 EST_HUMAN 22976 36196 0.46 0.0E+00 AV644766.1 EST_HUMAN 22977 36196 0.46 0.0E+00 BF438218.1 EST_HUMAN 23020 36274 2.5 0.0E+00 BF438218.1 EST_HUMAN 23021 36229 2.75 0.0E+00 BF6451742.1 EST_HUMAN 23020 36279 2.75 0.0E+00 BE642720.1 EST_HUMAN 23021 36307 1.9 0.0E+00 BE082720.1 EST_HUMAN 23081 <	10252	Ш	36110		0.0E+00	AV716271.1	EST_HUMAN	AV716271 DCB Homo saplens cDNA clone DCBBDC09 6'
22830 36143 2.36 0.0E+00 Al631618.1 EST_HUMAN 22830 36144 2.36 0.0E+00 Al631818.1 EST_HUMAN 22836 36149 0.49 0.0E+00 11545730 NT EST_HUMAN 22876 36196 0.46 0.0E+00 AU22428.1 EST_HUMAN 22876 36196 0.46 0.0E+00 AV64728.1 EST_HUMAN 22976 36214 2.5 0.0E+00 AV64766.1 EST_HUMAN 22966 36234 2.75 0.0E+00 AV64766.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE48278.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE649213.1 EST_HUMAN 23020 36276 0.75 0.0E+00 BE649213.1 EST_HUMAN 23020 36276 0.75 0.0E+00 BE649213.1 EST_HUMAN 23040 36306 1.9 0.0E+00 BE649213.1 EST_HUMAN 23040 36	10252		36111	0.55	0.0E+00	AV716271.1	EST_HUMAN	AV716271 DCB Homo sapiens cDNA clone DCBBDC09 5'
22930 36144 2.36 0.0E+00 AI631818.1 EST_HUMAN 22945 36149 0.49 0.0E+00 11545730 NT 22945 36149 0.49 0.0E+00 11545730 NT 22970 36190 0.64 0.0E+00 AU722429.1 EST_HUMAN 22965 36214 2.5 0.0E+00 BF485218.1 EST_HUMAN 22966 36231 2.75 0.0E+00 BF485218.1 EST_HUMAN 23020 36231 2.75 0.0E+00 BF485218.1 EST_HUMAN 23020 36232 2.75 0.0E+00 BE549213.1 EST_HUMAN 23021 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23020 36279 2.79 0.0E+00 BE649213.1 EST_HUMAN 23021 36336 0.5 0.0E+00 BE649273.1 EST_HUMAN 23049 36336 0.5 0.0E+00 BE042720.1 EST_HUMAN 23106 36336	10282		36143	2.36	0.0E+00	Al631818.1	EST_HUMAN	ws36e03.x1 NCI_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2300188 3' similer to TR:Q61204 Q61204 NOTCH2-LiKE;
22836 36149 0.49 0.0E+00 11545730 NT 22945 38159 1.52 0.0E+00 T03078.1 EST_HUMAN 22970 36190 0.64 0.0E+00 AU122429.1 EST_HUMAN 22976 36196 0.46 0.0E+00 BF436218.1 EST_HUMAN 22905 36234 2.5 0.0E+00 BF436218.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BF436213.1 EST_HUMAN 23035 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23040 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23041 36307 1.9 0.0E+00 BE649213.1 EST_HUMAN 23081 36279 2.79 0.0E+00 BE640270.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE640270.1 EST_HUMAN 23081 36308 1.3 0.0E+00 BE040270.1 EST_HUMAN 23100 36336 0.86<	10282		36144	2.36	0.0E+00	AI631818.1	EST HUMAN	ws36e03.x1 NCI_CGAP_Kid11 Homo seplens cDNA clone IMAGE:2300188 3' similar to TR:Q61204 Q61204 NOTCH2-LIKE;
22945 36159 1.52 0.0E+00 ITO3078.1 EST_HUMAN 22970 36190 0.64 0.0E+00 AU122429.1 EST_HUMAN 22976 36196 0.46 0.0E+00 BF436218.1 EST_HUMAN 22905 36234 2.5 0.0E+00 BF436218.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BF436213.1 EST_HUMAN 23045 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23050 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23060 36276 0.76 0.0E+00 BE649213.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE649213.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE6492213.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE6492213.1 EST_HUMAN 23081 36308 1.33 0.0E+00 BE6492213.1 EST_HUMAN 23106 <t< td=""><td>10288</td><td></td><td></td><td>0.49</td><td>0.0E+00</td><td>11545730</td><td>N</td><td>Homo saplens Gigaxonin (GAN), mRNA</td></t<>	10288			0.49	0.0E+00	11545730	N	Homo saplens Gigaxonin (GAN), mRNA
22970 36190 0.64 0.0E+00 AU122429.1 EST_HUMAN 22976 36196 0.46 0.0E+00 BF436218.1 EST_HUMAN 22966 36234 2.5 0.0E+00 BF436218.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BF486213.1 EST_HUMAN 23035 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23045 36276 0.75 0.0E+00 BE549213.1 EST_HUMAN 23060 36279 2.79 0.0E+00 BE649213.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE649213.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE649273.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE682720.1 EST_HUMAN 23081 36308 1.33 0.0E+00 BE082720.1 EST_HUMAN 23100 36336 0.36 0.0E+00 BE0176251.1 EST_HUMAN 23105 <td< td=""><td>10298</td><td></td><td>Н</td><td>1.52</td><td>0.0E+00</td><td></td><td>EST_HUMAN</td><td>FB23A4 Fetal brain, Stratagene Homo saplens cDNA clone FB23A4 3'end</td></td<>	10298		Н	1.52	0.0E+00		EST_HUMAN	FB23A4 Fetal brain, Stratagene Homo saplens cDNA clone FB23A4 3'end
22976 36196 0.46 0.0E+00 BF436218.1 EST_HUMAN 22965 36214 2.5 0.0E+00 BF436218.1 EST_HUMAN 22966 36231 2.75 0.0E+00 AV517960.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE649213.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE649213.1 EST_HUMAN 23040 36276 2.79 0.0E+00 BE649213.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE649272.1 EST_HUMAN 23081 36308 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36316 0.6E+00 BE082720.1 EST_HUMAN 23084 36316 0.6E+00 BE082720.1 EST_HUMAN 23107 1.33 0.0E+00 BE082720.1 EST_HUMAN 23108 36336 1.33 0.0E+00 BE017655.1 EST_HUMAN 23105 36336 1.33 0.0E+00	10323			0.64	0.0E+00		EST HUMAN	AU122429 MAMMA1 Homo sapiens cDNA clone MAMMA1002368 5'
22965 36214 2.5 0.0E+00 BF436218.1 EST_HUMAN 22906 36231 2.75 0.0E+00 AV617960.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE649213.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE649213.1 EST_HUMAN 23036 36276 2.79 0.0E+00 BE649213.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE649272.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE082720.1 EST_HUMAN 23082 36316 0.6E+00 BE082720.1 EST_HUMAN 23084 36325 0.8E 0.0E+00 BE082720.1 EST_HUMAN 23109 36331 1.33 0.0E+00 BE082720.1 EST_HUMAN 23105 36332 1.33 0.0E+00 BE043245.1 EST_HUMAN 23106 36336 2.49 0.0E+00 BE04325.1 EST_HUMAN 23166 36336 <td< td=""><td>10329</td><td>1</td><td>36196</td><td>0.46</td><td></td><td>5921</td><td>Į,</td><td>Homo septens triple functional domain (PTPRF interacting) (TRIO), mRNA</td></td<>	10329	1	36196	0.46		5921	Į,	Homo septens triple functional domain (PTPRF interacting) (TRIO), mRNA
23016 0.097 0.0E+00 AV664766.1 EST_HUMAN 23026 36236 2.75 0.0E+00 BE646213.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE646213.1 EST_HUMAN 23036 36276 0.0E+00 BE646213.1 EST_HUMAN 23081 36276 1.9 0.0E+00 BE781742.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE082720.1 EST_HUMAN 23082 36316 0.66 0.0E+00 BE082720.1 EST_HUMAN 23084 36325 0.86 0.0E+00 BE082720.1 EST_HUMAN 23109 36331 1.33 0.0E+00 BE082720.1 EST_HUMAN 23106 36332 1.33 0.0E+00 BE743216.1 EST_HUMAN 23107 36336 2.49 0.0E+00 BE647655.1 EST_HUMAN 23108 36336 0.57 0.0E+00 BE67765.1 EST_HUMAN 23166 36338 0.05+00	10348		36214	2.5	1		EST_HUMAN	nab45e12.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sapiens cDNA done IMAGE:32652713'
23015 36231 2.75 0.0E+00 AW517860.1 EST_HUMAN 23020 36236 8.82 0.0E+00 BE549213.1 EST_HUMAN 23036 36279 0.75 0.0E+00 BE743213.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE781742.1 EST_HUMAN 23084 36308 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36316 0.66 0.0E+00 BE082720.1 EST_HUMAN 23084 36325 0.86 0.0E+00 BE082720.1 EST_HUMAN 23109 36331 1.33 0.0E+00 BE743216.1 EST_HUMAN 23106 36332 2.49 0.0E+00 BE617655.1 EST_HUMAN 23107 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23164 36339 1.01 0.0E+00 BE617655.1 INT 23165 36339 1.01 0.0E+00 BE617655.1 INT 23165 36339	10349	- 1		0.97			EST_HUMAN	AV654765 GLC Homo saplens cDNA clone GLCDZC07 3'
23020 36236 8.82 0.0E+00 BE549213.1 EST HUMAN 23035 36251 0.75 0.0E+00 11436005 INT 23060 36270 2.79 0.0E+00 BE781742.1 EST HUMAN 23081 36307 1.9 0.0E+00 BE082720.1 EST HUMAN 23084 36308 1.9 0.0E+00 BE082720.1 EST HUMAN 23084 36316 0.66 0.0E+00 BE082720.1 EST HUMAN 23100 36331 1.33 0.0E+00 BE743216.1 EST HUMAN 23105 36332 1.33 0.0E+00 BE1742516.1 EST HUMAN 23105 36336 2.49 0.0E+00 BE617655.1 EST HUMAN 23164 36336 0.57 0.0E+00 BE617655.1 EST HUMAN 23165 36380 1.01 0.0E+00 BE617655.1 EST HUMAN 23165 36381 1.02 0.0E+00 BE6172264.1 EST HUMAN 23165 3638	10369	23015	36231	2.75	0.0E+00		EST HUMAN	xu74b01x1 NCI_CGAP_Kid8 Homo sapiens cDNA clone IMAGE:2807401 3' similar to gb:M89086 MOESIN (HUMAN);
23035 36251 0.75 0.0E+00 11436005 INT 23060 36279 2.79 0.0E+00 BE781742.1 EST_HUMAN 23081 36307 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36308 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36316 0.66 0.0E+00 PR0321 NT 23100 36332 0.0E+00 BE743215.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE1742515.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE1742515.1 EST_HUMAN 23105 36336 0.0E+00 BE1742515.1 EST_HUMAN 23167 36336 0.0E+00 BE17455.1 EST_HUMAN 23167 36336 0.0E+00 BE172264.1 NT 23168 363392 0.0E+00 BE172254.1 EST_HUMAN 23165 363392 0.02 0.0E+00 BE172254.1 EST_HUMAN <tr< td=""><td>10374</td><td>23020</td><td>36236</td><td>8.82</td><td>0.0E+00</td><td>Γ</td><td>EST HUMAN</td><td>601078764F1 NIH MGC_12 Homo saplens cDNA clone IMAGE:3484703 6</td></tr<>	10374	23020	36236	8.82	0.0E+00	Γ	EST HUMAN	601078764F1 NIH MGC_12 Homo saplens cDNA clone IMAGE:3484703 6
23060 3627b 2.79 0.0E+00 BE781742.1 EST HUMAN 23081 36307 1.9 0.0E+00 BE082720.1 EST HUMAN 23084 36308 1.9 0.0E+00 BE082720.1 EST HUMAN 23084 36316 0.66 0.0E+00 P08032.1 NT 23100 36332 0.86 0.0E+00 BE743216.1 EST HUMAN 23105 36336 2.49 0.0E+00 BE743216.1 EST HUMAN 23105 36336 2.49 0.0E+00 BE017655.1 EST HUMAN 23144 36336 0.57 0.0E+00 BE017655.1 EST HUMAN 23167 36336 0.57 0.0E+00 BE017655.1 EST HUMAN 23167 36336 0.45 0.0E+00 BE017655.1 EST HUMAN 23168 36339 1.01 0.0E+00 BE017256.1 RST HUMAN 23169 36339 1.02 0.0E+00 BE172254.1 EST HUMAN 23228 36463 </td <td>10389</td> <td>23035</td> <td>36251</td> <td>0.75</td> <td>0.0E+00</td> <td>3005</td> <td>Z</td> <td>Homo saplens hypothetical protein DKFZp761P1010 (DKFZp761P1010), mRNA</td>	10389	23035	36251	0.75	0.0E+00	3005	Z	Homo saplens hypothetical protein DKFZp761P1010 (DKFZp761P1010), mRNA
23081 36307 1.9 0.0E+00 BE082720.1 EST_HUMAN 23088 36308 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36316 0.66 0.0E+00 Y08032.1 NT 23100 36325 0.86 0.0E+00 BE743216.1 EST_HUMAN 23100 36336 2.49 0.0E+00 BE743216.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23127 36356 0.57 0.0E+00 BE617655.1 EST_HUMAN 23127 36356 0.57 0.0E+00 BE617655.1 EST_HUMAN 23164 36380 1.01 0.0E+00 BF017655.1 EST_HUMAN 23165 36381 1.02 0.0E+00 BF017256.1 RST_HUMAN 23165 36382 0.45 0.0E+00 BF017256.1 RST_HUMAN 23165 36382 1.02 0.0E+00 BE172256.1 EST_HUMAN 23228 36463 </td <td>10414</td> <td>i</td> <td>36279</td> <td>2.79</td> <td>0.0E+00</td> <td></td> <td>EST_HUMAN</td> <td>801487418F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870700 5'</td>	10414	i	36279	2.79	0.0E+00		EST_HUMAN	801487418F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3870700 5'
23081 36308 1.9 0.0E+00 BE082720.1 EST_HUMAN 23084 36316 0.66 0.0E+00 Y08032.1 NT_HUMAN 23100 36325 0.86 0.0E+00 BE743216.1 EST_HUMAN 23100 36336 2.49 0.0E+00 BE743216.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23145 36336 0.57 0.0E+00 BE617655.1 EST_HUMAN 23147 36336 0.57 0.0E+00 BE617655.1 EST_HUMAN 23148 36336 0.57 0.0E+00 BF617655.1 EST_HUMAN 23156 36331 1.01 0.0E+00 BF617256.1 EST_HUMAN 23165 36332 1.02 0.0E+00 BF617226.1 EST_HUMAN 23165 36332 1.02 0.0E+00 BF172264.1 EST_HUMAN 23165 36392 1.02 0.0E+00 BE172254.1 EST_HUMAN 23228	10435	- 1	36307	1.9	0.0E+00		EST_HUMAN	RC2-BT0642-150200-012-d03 BT0642 Homo saplens cDNA
2308B 36316 0.66 0.0E+00 Y08032.1 NT 23094 36325 0.86 0.0E+00 Al656890.1 EST_HUMAN 23100 36331 1.33 0.0E+00 BE743216.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE743215.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE017655.1 EST_HUMAN 23127 36336 0.57 0.0E+00 BE017655.1 EST_HUMAN 23146 36380 1.01 0.0E+00 BF01755.1 INT 23166 36382 0.45 0.0E+00 BF017254.1 INT 23167 36382 1.02 0.0E+00 BF172254.1 EST_HUMAN 23165 36382 1.02 0.0E+00 BE172254.1 EST_HUMAN 23228 36463 2.76 0.0E+00 BF172254.1 EST_HUMAN	10435	- 1	36308	1.9	0.0E+00		<u>ا</u> ا	RC2-BT0642-150200-012-d03 BT0642 Homo saplens cDNA
23024 363.25 0.86 0.0E+00 Al656890.1 EST_HUMAN 23100 36331 1.33 0.0E+00 BE743216.1 EST_HUMAN 23100 36332 1.33 0.0E+00 BE617655.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23127 36336 0.57 0.0E+00 BE617655.1 EST_HUMAN 23167 36386 0.57 0.0E+00 BR905.1 EST_HUMAN 23168 36382 0.45 0.0E+00 B797254.1 NT 23169 36382 0.45 0.0E+00 BF172264.1 EST_HUMAN 23165 36382 1.02 0.0E+00 BE172254.1 EST_HUMAN 23165 36382 1.02 0.0E+00 BE172254.1 EST_HUMAN 23228 36463 2.76 0.0E+00 BC172254.1 EST_HUMAN	10442	ı	36316	0.66	0.0E+00		Ę	Human endogenous retrovirus-K, LTR U5 and gag gene
23100 36331 1,33 0.0E+00 BE743216.1 EST_HUMAN 23100 36332 1,33 0.0E+00 BE743216.1 EST_HUMAN 23105 36336 2,49 0.0E+00 BE617655.1 EST_HUMAN 23127 36356 0.57 0.0E+00 BE617655.1 EST_HUMAN 23164 36380 1.01 0.0E+00 BR905.1 EST_HUMAN 23165 36382 0.01 0.0E+00 BR9765.1 BST_HUMAN 23166 36382 0.04 0.0E+00 BR9765.1 BST_HUMAN 23167 36382 0.04 0.0E+00 BR9765.1 BST_HUMAN 23167 36382 1.02 0.0E+00 BE172264.1 BST_HUMAN 23168 36383 1.02 0.0E+00 BE172254.1 EST_HUMAN 23228 36463 2.76 0.0E+00 BC172254.1 EST_HUMAN	10448	ı	36325	0.86	0.0E+00		EST_HUMAN	tt64e07.x1 NCI_CGAP_GC6 Homo septens cDNA clone IMAGE:2244612.3'
23100 36332 1.33 0.0E+00 BE743215.1 EST_HUMAN 23105 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23127 36356 0.57 0.0E+00 BE617655.1 EST_HUMAN 23154 36380 1.01 0.0E+00 B7675.1 INT 23156 36382 0.45 0.0E+00 B7675.1 INT 23165 36392 1.02 0.0E+00 BE172264.1 INT 23165 36393 1.02 0.0E+00 BE172264.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10454	- 1	36331	1.33	0.0E+00		EST_HUMAN	601573895F1 NIH_MGC_9 Homo sepiens cDNA clone IMAGE:3835198 5'
23105 38336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23105 38336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23127 38356 0.57 0.0E+00 H39805.1 EST_HUMAN 23154 36380 1.01 0.0E+00 D87675.1 NT 23165 36392 0.45 0.0E+00 D87681364.1 NT 23165 36392 1.02 0.0E+00 BE172264.1 EST_HUMAN 23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10454	- 1	36332	1.33	0.0E+00		EST_HUMAN	601573895F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3835198 6'
23105 36336 2.49 0.0E+00 BE617655.1 EST_HUMAN 23127 36356 0.57 0.0E+00 H39805.1 EST_HUMAN 23154 36380 1.01 0.0E+00 D87675.1 NT 23156 36392 0.45 0.0E+00 AF081364.1 NT 23165 36393 1.02 0.0E+00 BE172264.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10459	. 1	36336	2.49	0.0E+00		EST_HUMAN	601441723T1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3845956 3'
23127 36355 0.57 0.0E+00 H39805.1 EST_HUMAN 23154 36380 1.01 0.0E+00 D87675.1 NT 23156 36382 0.45 0.0E+00 AF081364.1 NT 23165 36392 1.02 0.0E+00 BE172264.1 EST_HUMAN 23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10459	23105	36336	2.49	0.0E+00		EST_HUMAN	801441723T1 NIH_MGC_65 Homo saplens cDNA clone IMAGE:3846968 3'
23154 36380 1.01 0.0E+00 D87675.1 NT 23156 36382 0.45 0.0E+00 AF081364.1 NT 23165 36392 1.02 0.0E+00 BE172264.1 EST_HUMAN 23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10481	23127	36355	0.57	0.0E+00		EST_HUMAN	yp01a10.r1 Soares breast 3NbHBst Homo sepiens cDNA clone IMAGE:186138 6'
23156 36382 0.45 0.0E+00 AF081364.1 NT 23165 36392 1.02 0.0E+00 BE172264.1 EST_HUMAN 23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10508	23154	36380	1.01	0.0E+00		LN L	Homo saplens DNA for amyloid precursor protein, complete cds
23165 36392 1.02 0.0E+00 BE172264.1 EST_HUMAN 23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10510	23156	36382	0.45	0.0E+00	1.1	TN	Synthetic construct CD30 ligand-exotoxin A fusion protein (CD30L-ETA fusion) mRNA, partial cds
23165 36393 1.02 0.0E+00 BE172254.1 EST_HUMAN 23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10519	23165	36392	1.02	0.0E+00		EST_HUMAN	MR0-HT0569-270300-006-e12 HT0569 Homo sepiens cDNA
23229 36463 2.76 0.0E+00 AV711075.1 EST_HUMAN	10519	23165	36393	1.02	0.0E+00		EST_HUMAN	MR0-HT0569-270300-008-e12 HT0569 Homo saplens cDNA
1	10532	23220	36463	2.76	0.0E+00		EST_HUMAN	AV711075 Cu Homo sapiens cDNA clone CuAAKG05 5'

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					1		
Probe SEQ ID NO:	Econ SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acession No.	Top Hit Database Source	Top Hit Descriptor
10532	23229	36464	2.76	0.0E+00	AV711075.1		AV711075 Cu Homo sepiens cDNA clone CuAAKG05 5'
10534	23231		2.13	0.0E+00	П	EST_HUMAN	RC3-ST0197-120200-015-e03 ST0197 Homo saplens cDNA
10542	23238	36472	7.02	0.0E+00	AW963563.1	EST_HUMAN	EST375636 MAGE resequences, MAGH Homo sapiens cDNA
10555	23251	36487		0.0E+00	11431124 NT		Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABC43), mRNA
10555	23251	36488	3.19	0.0E+00	11431124 NT		Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3), mRNA
10559	23255	38492	2.09	0.0E+00	AW057621.1	EST_HUMAN	wy61f09.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sepiens cDNA clone IMAGE:2553055 3' similar to TR:Q60566 Q60566 VDX;
10567	23262	36499		0.0E+00	BE243270.1	EST_HUMAN	TCAAP3D0917 Pedietric acute myelogenous leukemia celi (FAB M1) Baylor-HGSC project=1CAA Hamo saplens cDNA clone TCAAP0917
10568	23263	36500	2.85	0.0E+00	A1652239.1	EST_HUMAN	wb28a12x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:2306974 3' similar to contains element MSR1 MSR1 repetitive element;
10568	23263	36501		0.0E+00	A1662239.1	EST_HUMAN	wb28a12x1 NCI_CGAP_GC9 Homo sapiens cDNA clone IMAGE:2306974 3' sImilar to contains element MSR1 MSR1 repetitive element;
10573	23268	36508		0.0E+00	BF306642.1	EST_HUMAN	601888704F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4122649 5'
10580	23275	36512	5.06	0.0E+00	11545911 NT	TN	Homo sapiens NOD2 protein (NOD2), mRNA
10580	23275	36513	5.08	0.0E+00	11545911 NT	NT	Homo sapiens NOD2 protein (NOD2), mRNA
10596	23290	36528	1.98	0.0E+00	AW 404795.1	EST_HUMAN	UIHF-BLO-ecm-d-04-0-UIM NIH MGC 37 Homo sapiens cunk ciane invalue: 3039363 3
10600	23294	36533	3.17		11424829 NT	N	Homo saplens hypothetical protein FLJ20079 (FLJ20079), mKNA
10801	23295				4504536 NT	LN.	Homo sapiens 5-hydroxydyptamine (serotonin) receptor 1E (n i R IE) mixiva
10601	23295			0.0E+00	4504536 NT	LN	Homo sapiens 5-nydroxydypuanine (secondini) reception in (in think) mixty mixty (in the condining mixty) in the condining mixty of the co
10602	23296			0.0E+00	Al991827.1	EST_HUMAN	WISZDOSZI SOBJES DIECKGIRGIE COUNTINO PARIO CON COUNTINO CENTRALINA CENTRALIN
10605	23299			0.0E+00		EST_HUMAN	601505204FZ NIH MGC_ / I Homo septems count circums invoce. 3500000 5
10609	23303				BE89163	EST_HUMAN	60143452271 NIT MGC_/Z HOURD SQUARIS COINT CORE INTO CENSOR OF THE CONTROL OF THE
10612	23306					Ž!	Tomo saprens myosh, neavy pulypopude z, exercial musch, and (mm. z.) mw
10612	23306					Z	Tomo sapiens myosni, neavy pulypopules z, skereta musous, akun (m. 1.2), m. 57.
10619	23312	36551	1.4			LN	Homo sapiens mkny for no protein, partar
10619	23312	36552	1.4		AB014608.1	Ŋ	Homo saplens mRNA for KIAA0708 protein, perital cds
10628	23321	36559	1.31	0.0E+00	BE903304.1	EST_HUMAN	601674332F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3957343 5
		L		<u> </u>		1444	zp95b11.r1 Stratagene muscle 937209 Homo sapiens cDNA clone IMAGE:627933 5' similar to gb:X03740 AXOCIN DEAMY CHAIN SKELETAL MISCLE (HLIMAN):
10631				0.0=+00	AA190900.1	NAME TO THE	MICOLEMENT OF THE PROPERTY OF
10652	23343			0.0E+00		ESI HOMAN	COLOCOCAZET INITIAMOS TO Home capiene chiNA clone IMAGE:38325755
10660	23351			1		ESI HUMAN	CO 1302504FT INIT MICE AN INC. 20 Home contant china IMAGE:3832575 57
10660	23351			1	BE729706.1	ESI HUMAN	001302004FT NIT MOULE CONTINUE SEPTEMBER CONTINUES FOR CON
10661	23352	36590	33.99	0.0E+00	AV727362.1	EST HOMAN	AV Zrocz nie nomo sapienie centa ciule i i energia

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Table 4
Single Exon Probes Expressed in

			gb:M60854 40S		llar to contains	llar to contains	lar to contains						:X17118 IG MU	3,	3,														
Single Exon Probes Expressed in Brain	Top Hit Descriptor	AV727362 HTC Homo saplens cDNA clone HTCAQH08 5'	xy04g10.x1 NCI_CGAP_Lym12 Homo sapiens cDNA clone IMAGE:2882226 3' similar to gb:M60864 40S RIBOSOMAL PROTEIN S16 (HUMAN);	AU135741 PLACE1 Homo sapiens cDNA clone PLACE1002784 6'	hg13d02.x1 Soares, NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2946476 3' similar to contains element MSR1 repetitive element;	hg13d02.x1 Scares, NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2845475.3' similar to contains element MSR1 repetitive element;	hg13d02.x1 Spares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2846475 3' similar to contains element MSR1 repetitive element :	H.saplens mRNA for H1 histamine receptor	HSC3IC031 normalized infant brain cDNA Homo saplens cDNA clone c-3ic03	Homo sapiens RGH1 gene, retrovirus-like element	Homo saplens EphA7 (EPHA7) mRNA	Homo saplens EphA7 (EPHA7) mRNA	xw66f01x1 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2832985 3' similar to gb:X17115 IG MU CHAIN C RECION (HUMAN);	UI-H-Bi3-alh-a-01-0-UI,s1 NCI CGAP Sub5 Homo sapiens cDNA clone IMAGE:2738649 3	UI-H-BI3-alh-a-01-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2736649.3	Homo sapiens ribosomal protein L31 (RPL31) mRNA	Homo sapiens mRNA for KIAA0667 protein, partial cds	601119248F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3029219 5'	Homo saplens mRNA for KIAA0545 protein, partial cds	AU124106 NT2RM2 Homo sapiens cDNA clone NT2RM2001676 5	Homo saplens mRNA for KIAA1117 protein, partial cds	Homo saplens mRNA for KIAA1117 protein, partial cds	801582046F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3936539 6	602141405F1 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4302432 5	801186342F1 NIH_MGC_8 Homo sapiens cDNA clone IMAGE:3544259 5'	AU118386 HEMBA1 Homo sapiens cDNA clone HEMBA1003486 5'	qf43c03.x1 Scares_testis_NHT Homo sapiens cDNA clone IMAGE:1762772 3'	qf43c03.x1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:1762772 3'	QV4-ST0234-121199-032-b06 ST0234 Homo sapiens cDNA
gle Exon Pro	Top Hit Database Source	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	EST HUMAN	NT	님	١	EST_HUMAN	EST HUMAN	EST_HUMAN	ΙΝ	L	EST_HUMAN	NT	EST_HUMAN	Ę	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
Sin	Top Hit Acession No.	-00 AV727362.1	-00 AW 516055.1	00 AU135741.1	.00 AW 69333.1	00 AW59333.1	00 AW 69333.1		Γ		4758281 NT	4758281 NT	00 AW338094.1	0.0E+00 AW461230.1	0.0E+00 AW 451230.1	4506632 NT	0.0E+00 AB014567.1												00 AW391937.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 Z34897.1	0.0E+00 F13069.1	0.0E+00 D10083.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00 A	0.0E+00	0.0E+00 AB029040.1	0.0E+00 AB029040.1	0.0E+00 BE792155.1	0.0E+00 BF684061.1	0.0E+00 BE269288.1	0.0E+00 AU118386.1	0.0E+00 A	0.0E+00 A	0.0E+00 A
	Expression Signal	33.99	9.59	3.18	3.41	3.41	3.41	1.89	2.97	1.79	1.33	1.33	2.13	4.62	4.62	11.67	2.53	1.98	2.04	1.71	1.45	1.45	4.04	59.14	1.3	6.9	6.53	6.53	3.04
	ORF SEQ ID NO:	36591	36608	36613	36617	36618	36619	36620	36621	36629	36834	36635	36648	36849	36650		36662	36670	36683		36697	36698	36702		36703	38706	36710	38711	36712
	Exan SEQ ID NO:	23352		23371	23377	23377	23377	23379	23380	23391	23396	23396	23407	23408	23408	13021	23411	23425	23439	23447	23454	23454	23459	23460	23461	23484	23469	23469	23470
	Probe SEQ ID NO:	10661	10674	10680	10686	10886	10686	10688	10689	10700	10708	10708	10718	10719	10719	10721	10723	10738	10754	10763	10771	10771	10778	10777	10778	10781	10786	10786	10787

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Probe SEQ ID (9	SEQ ID	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Acesslon No.	Top Hit Database Source	Top Hit Descriptor
10.798	23481	36721	4.39	0.0E+00	AF223391.1	N.	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced
							Homo saplens celcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively
10798	23481	36722		0.0E+00	AF2Z3391.1	Z	Spiroto Homo caniane Insulla recentor (INSR) mRNA
10807	23498	36726	9.57	0.01-00	11424720 N I	EST HIMAN	Profile Septeds insulin receptor (1909), Illinois QVc-UM0083-170400-191-406 UM0093 Homo septens cDNA
10814	23497	36734		0.00	AW804516.1		QVo-UM0093-170400-191-d06 UM0093 Hamo saplens cDNA
10815	23498	36735			BF340308.1	EST_HUMAN	602037014F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4184979 5'
10817	23500		52.94	0.0E+00	BE261209.1	EST HUMAN	601148357F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3163310 5
10821	23504	36743	2.37	0.0E+00	AB029040.1	NT	Homo sapiens mRNA for KIAA1117 protein, partial cds
10824	23507	36746	1.69	0.0E+00	AB007932.1	NT	Homo sapiens mRNA for KIAA0463 protein, partial cds
10828	23510		3.47	0.0E+00	U50326.1	NT	Human protein kinase C substrate 80K-H (PRKCSH) gene, exon 15-17
10832	23514			0.0E+00	BE773036.1	EST_HUMAN	RC1-FT0134-170700-012-f07 FT0134 Homo saplens cDNA
10832	23614	36756	1.55		BE773036.1	EST_HUMAN	RC1-FT0134-170700-012-f07 FT0134 Homo saplens cDNA
10838	23520	36762		0.0E+00	W21826.1	EST_HUMAN	57E10 Human retina cDNA Tsp509I-cleaved sublibrary Homo saplens cDNA not directional
							ob32e07.s1 NCI_CGAP_Kid5 Homo sapiens cDNA clone IMAGE:1325412.3' similar to contains element
10854	23534	36779	136.91	0.0E+00	AA740782.1	EST_HUMAN	MSR1 repetitive element;
10857	23537	36783	2.05	0.0E+00	AW 466922.1	EST_HUMAN	ha04h04.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:28/2/393
10863	23643	36790		0.0E+00	AF252303.1	٦ ا	Homo sapiens signaling lymphocytic activation molecule (SLAM) gene, expl. 2011
10879	23559	36806	7.34	0.0E+00	C05089.1	EST_HUMAN	C05089 Human heart cDNA (YNakamura) Homo sapiens cDNA clone 3NHC4817
10886	23566	36814	2.31	0.0E+00	AA746376.1	EST_HUMAN	oa56h01,r1 NCI_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1309009 5
10886	23566	36815	2.31	0.0E+00	AA746375.1	EST_HUMAN	oa56h01.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1309009 5
10895	23575			0.0E+0	JM78448.1	EST_HUMAN	EST00596 Fetal brain, Stratagene (cat#336206) Homo sapiens cDNA clone HFBCC26
10895	23575	36826	3.74	0.0E+00	M78448.1	EST_HUMAN	EST00596 Fetal brain, Stratagene (cat#836206) Homo sapiens cUNA clone HFBCC26
10898	23578	36827	6.82	0.0E+00	AL157608.1	EST_HUMAN	DKFZp761J2116_r1 761 (synonym: hamy2) Homo sapiens cDNA cione DKr zp761J2116 5
10910	23590	36836	5.81	0.0E+00	AU116988.1	EST_HUMAN	AU116988 HEMBA1 Homo sapiens cDNA clone HEMBA1000424 5
10924	23604		1.64	0.0E+00	AV693656.1	EST_HUMAN	AV693656 GKC Homo saplens cDNA clone GKCCNC03 5
10932	23612		2.09	0.0E+00	BF366553.1	EST_HUMAN	IL3-NT0104-200500-143-A07 NT0104 Homo saplens cDNA
10955	18399	31311	2.73	0.0E+00	AB035266.1	TN	Homo sapiens mRNA for neurexin II, complete cds
10955	18399	31312	2.73	0.0E+00	AB035266.1	ΝΤ	Homo sapiens mRNA for neurexin II, complete cds
10960	23636	36887	2.64	0.0E+00	BE182360.1	EST_HUMAN	PMO-HT0845-060500-002-E05 HT0645 Homo sapiens cDNA
10960	23636	36888	2.64	0.0E+00	BE182360.1	EST HUMAN	PM0-HT0645-060500-002-E05 HT0645 Homo sapiens cUNA
10961	23637		1.4		0.0E+00 AV701152.1	EST HUMAN	AV701152 ADA Homo sapiens cDNA clone ADAAADU6 5
10980	23655	36908	4.07		0.0E+00 BE896423.1	EST_HUMAN	601439092F1 NIH_MGC_72 Homo sapiens CUNA cione IMAGE:3824142 3

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					JIO .	igle Exon Pro	Single Exon Probes Expressed in Brain
Probe SEQ ID NO:	Exen SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E	Top Hit Acession	Top Hit Database Source	Top Hit Descriptor
10989	23663	36919		L	AW 500307 1	FST HIMAN	HEBNO and ADSO III A NIU MSC SA Lambouries - ANA ANA
10989			1.85		0.0E+00 AW500307.1	EST HUMAN	U-HF-BN0-ako-d-02-0-01 r1 NIH MGC 50 Home septems contact and IMA CE-20177019 5
10000	aaacc	00000					bb78c04.y1 NIH_MGC_10 Homo saplens cDNA done IMAGE:3048488 5' similar to gb:y00346_cdo1 POLYADENYLATE-BINDING PROTEIN (HUMAN); gb:X85563 M.musculus mRNA for poly(A) binding
11016	23688	36949	1.77	0.0E+	0.0E+00 BE018293.1	EST HUMAN	protein (MOUSE);
11016	23688	36950	1.71	0.0E+00	0.0E+00 BF528907 1	EST HUMAN	802043377F1 NCT CGAP Rings Home seniors CDNA close IMAGE: 4181083 5
11016	23688	36951	1.77	0.0E+00	0.0E+00 BF528907.1	EST HUMAN	802043377F1 NCI CGAP RmR7 Home semiens cDNA close 1446 CE 4484782 51
11028	25133	36964	1.27	0.0E+00	0.0E+00 AW387786.1	EST HUMAN	MRA-ST0118-041099-010-A12 ST0118 Homo septems cDNA
11028	25133	36965	1.27	0.0E+00	0.0E+00 AW387768.1	EST HUMAN	MR4-ST0118-041099-010-A12 ST0118 Homo septiens cDNA
11034	23705	36973	1.63	0.0E+00	4758281 NT	N L	Homo saplens EphA7 (EPHA7) mRNA
11035	23706	36974	8.73	0.0E+00	00 BE897953.1	EST HUMAN	801440448F1 NIH MGC 72 Homo saplens cDNA clone IMAGE 3025403 F
11037	23708	36977	1.89	0.0E+00	0.0E+00 Al458545.1	EST_HUMAN	ao88g11.x1 Schiller meningioma Homo sapiens oDNA done IMAGE-1952804.3
11037	23708	36978	1.89	0.0E+00		EST HUMAN	ao86g11.x1 Schiller meningiorna Homo saplens cDNA clone IMAGE:1952804.3
11051	23721	36992	2.78	0.0E+00		EST HUMAN	DKFZp434L0120 r1 434 (swngnym: htes3) Homo saplens oDNA clone DKFZp434L0120 r1
11083	23753	37028	1.61	0.0E+00	10880982	NT	Homo saplens gephyrin (GPH), mRNA
11085	23765	37031	3.98	0.0E+00	4758827 NT	LN	Home saplens neuredn III (NRXN3) mRNA
11086	23756	37032	2.67	0.0E+00	BF206561.1	EST HUMAN	601870902F1 NIH MGC 19 Homo septems cDNA clone IMAGE: 4104433 F.
11091	23761	37036	12.22	0.0E+00		EST HUMAN	UI-H-BI2-age-h-01-0-UI.s1 NCI CGAP Sub4 Homo septens aDNA clone INACE: 2724342 31
11095	23766	37040	4.23	0.0E+00		FN	Homo saplens mRNA for KIAA0717 protein, partial cds
11096	23766	37041	4.23	0.0E+00	0.0E+00 AB018260.1	FZ	Homo sapiens mRNA for KIAA0717 protein, partial cds
11098	23768	37043	2.69	0.0E+00	0.0E+00 BE206846.1	EST_HUMAN	be04d07.y1 NIH MGC 7 Homo sapiens cDNA done IMAGE:2823373 5' similar to TR:O76022 O76022 E1B- 55KDA-ASSOCIATED PROTEIN
11098	23768	37044	2.69	0.0E+00	0.0E+00 BE206846.1	EST HUMAN	be04407.y1 NIH_MGC_7 Homo seplens cDNA clone IMAGE:2823373 5' similar to TR:076022 076022 E1B-55KDA-ASSOCIATED PROTEIN
11110	23780	37056	1.9	0.0E+00	11526409	N	Homo sapiens KIAA0426 gene product (KIAA0426) mRNA
11124	23793	37069	1.62	0.0E+00	11024711 NT	LN L	Homo saplens myosin, heavy bolypeptide 4, skeletal muscle (MYHA) mRNIA
11127	20052	33133	1.5	0.0E+00	1,32832.1	NT	Homo saplens zinc finger homeodomain protein (ATBF1-A) mRNA complete ode
11131	23799	37074	3.84	0.0E+00	0.0E+00 BE148076.1	EST_HUMAN	RC3-HT0230-040500-110-h04 HT0230 Homo sapiens cDNA
11131	23799	37075	3.84	0.0E+00	0.0E+00 BE148076.1	EST_HUMAN	RC3-HT0230-040500-110-h04 HT0230 Homo sapiens cDNA
11154	23821	37101	1.96	0.0E+00	0.0E+00 AW673469.1	EST_HUMAN	be54d08.y3 NIH_MGC_10 Homo saplens cDNA done IMAGE:2900387 5' similar to TR:080275 060275 KIAA0522 PROTEIN ;
11154	23821	37102	1.96	0.0E+00	0.0E+00 AW673469.1	EST_HUMAN	ba54408.y3 NIH_MGC_10 Homo sapiens cDNA done IMAGE:2800367 5' similar to TR:O80275 O80275 KIAA0522 PROTEIN;

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	1.5		_		_			_						_					_		_	_	_	_			
Top Hit Descriptor	De04d07.71 NIH_MGC_7 Horne septens cDNA clone IMAGE:2823373 5' similar to TR:076022 076022 E1B 55KDA-ASSOCIATED PROTEIN	De04407.71 NIH_MGC_7 Home septens cDNA clone IMAGE:2823373 5' similar to TR:076022 076022 E1B 55KDA-ASSOCIATED PROTEIN	QV0-CT0225-101299-071-f06 CT0225 Homo saplens cDNA	nI42c08.s1 NCI_CGAP_Pr4 Homo saplens cDNA done IMAGE:1043342 similar to gb:M96178 ALPHA-ACTININ 1, CYTOSKELETAL ISOFORM (HUMAN):	wp06g08.x1 NCI CGAP Kid12 Homo sapiens cDNA clone IMAGE:2484004 3	drozbo8x1 NIH_MGC_3 Homo saplens oDNA clone IMAGE:2846919 5	UI-H-BW0-aij-d-07-0-UI.s1 NCI CGAP Sub6 Homo sapiens cDNA clone IMAGE-2729509 31	Homo saplens neurexin III (NRXN3) mRNA	601113903F1 NIH MGC 16 Homo sapiens cDNA clone IMAGE:33548nn 6'	601659088R1 NIH MGC 70 Homo sapiens aDNA clone IMAGE:3895018 3	601659088R1 NIH_MGC_70 Hamo saplens cDNA clone IMAGE:3895916 3'	IL5-HT0731-020500-077-f05 HT0731 Homo sapiens cDNA	UI-H-BW1-amv-a-05-0-UI.s1 NCI CGAP Sub7 Homo sanians cDNA clone IMA GE-2074121 2	DKFZp434G178 r1 434 (syronym: https3) Homo saplens cDNA clone DKFZp434G178 F	DKFZp434G178_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434G178 6'	wn83g03.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2452468 3' similar to gb:S37431 LAMININ RECEPTOR (HUMAN);	nz11c07.s1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:1287468 3' sImilar to TR:Q13686 Q13688 ALKB HOMOLOG PROTEIN .	nz11c07.s1 NCI_CGAP_GCB1 Homo septens oDNA clone IMAGE:1287468 3' similar to TR:Q13686 Q13686 ALKB HOMOLOG PROTFIN	601501090F1 NIH MGC 70 Homo sapiens cDNA clane IMAGE:3902028 5'	72712.x1 NCL CGAP_CLL1 Homo sapiens cDNA clone IMAGE:3295919 3' similar to TR:000409 000409 CHECKPOINT SUPPRESSOR 1	60/279335F1 NIH MGC 39 Homo sanians cDNA Alma MAGE 3841444 E1	601278335F1 NIH MGC 39 Homo saplens cDNA clone IMAGE:3811144 5	AV757420 BM Homo sapiens cDNA clone BMFAGH03 5'	Human endogenous retrovirus type K (HERV-K), aag, pol and env genes	Homo sapiens polycystic kidney disease-associated protein (PKD1) gene, complete cds	Homo sapiens polycystic kidney disease-associated protein (PKD1) gene, complete cds	AU138211 PLACE1 Homo saplens cDNA clone PLACE1008077 5'
Top Hit Database Source	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	L	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	N-	ΝT		EST_HUMAN
Top Hit Acession No.	BE206846.1	BE206846.1	AW753028.1	AA558707.1	AI934954.1	4W327895.1	4W292776.1	4758827	3E254058.1	3E965909.2	3E965909.2	3E185656.1	3F513960.1	1,046540.1	\L046540.1	1923116.1	A760913.1	A760913.1	E910546.1				Γ				00/AU138211.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00/	0.0E+00 /	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/Y	0.0E+00	0.0E+00 L	0.0E+00
Expression Signal	4.02	4.02	3.88	3.06	3.84	7.45	1.89	2.2	1.73	1.74	1.74	4.52	1.29	7.81	7.81	5.89	3.42	3.42	<u>2</u> .	7.9	1.79	1.79	1.61	1.52	10.31	10.31	4.69
ORF SEQ ID NO:	37321	37322	37324		į										37295	37306	37353	37354	37360	36413	36416	36417	36424	37367	37370	37371	37387
		24018		24025	18000	24026	25134	23173	23975	23978	23978	23979	23980	23993	23993	24003	24050	24050	24055	23183	23186	23186	23193	24061	24064	24084	24077
Probe SEQ (D NO:	11327	11327	11329	11334	11335	11338	11356	11362	11368	11371	11371	11372	11373	11387	11387	11397	11401	11401	11406	11416	11419	11419	11428	11457	11481	11461	11476
	SEQ ID ORF SEQ Signal (Top) Hit Top Hit Acession NO: Signal BLASTE No. Source	Exon ORF SEQ Expression ID NO: Crop) Hit Signal Top Hit Acession Signal Top Hit Top Hit Acession No. Top Hit Acession Source Source Top Hit Acession Source Source Source Detablese Source Source Source Source	Exon SEQ ID ID NO: ORF SEQ Signal Expression RLAST E Value (Top Hit No. Top Hit Source Value Top Hit No. Top Hit Source Source 24018 37321 4.02 0.0E+00 BE206846.1 EST_HUMAN 24018 37322 4.02 0.0E+00 BE206846.1 EST_HUMAN	Exon NO: ORF SEQ ID NO: Expression Signal Most Similar (Top) Hit Nelson Top Hit Acession No. Top Hit Source Source 24018 37322 4.02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 3.88 0.0E+00 AW753028.1 EST_HUMAN	Exon No: ORF SEQ ID NO: Expression Signal (Top) Hit BLAST E Velue Top Hit No. Top Hit Source Source 24018 37321 4.02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 3.88 0.0E+00 AM753028.1 EST_HUMAN 24025 3.08 0.0E+00 AM553028.1 EST_HUMAN EST_HUMAN EST_HUMAN EST_HUMAN	Exon No: ORF SEQ ID NO: Expression Signal (Top) Hit HLAST E Velue Top Hit No. Top Hit Source Source 24018 37321 4.02 0.05+00 BE206846.1 EST_HUMAN 24026 37324 3.88 0.05+00 AM753028.1 EST_HUMAN 24026 3.06 0.05+00 AM558707.1 EST_HUMAN 18000 30623 3.84 0.05+00 AM588707.1 EST_HUMAN	Exon No: ORF SEQ ID ID NO: Expression Signal (Top) Hit Acession Signal Top Hit Acession Source Noil Noil Signal Top Hit Acession Source Noil Noil Source Source Noil Noil Source Noil Noil Noil Noil Noil Noil Noil Noil	Exon No: ORF SEQ ID ID NO: Expression Signal (Top) Hit Acession Signal Top Hit Acession Source Noil BLAST E No. Top Hit Acession Source Source Noil BLAST E Noil BLAST E Noil BLAST E Noil BLAST E Source Noil Wellue 24018 37322 4.02 0.05+00 BE206846.1 EST_HUMAN 24026 37324 3.88 0.05+00 AW753028.1 EST_HUMAN 24026 3.06 0.05+00 AW753028.1 EST_HUMAN 24026 3.08 0.05+00 AW327895.1 EST_HUMAN 24026 3.7330 7.46 0.05+00 AW327895.1 EST_HUMAN 24026 37330 7.46 0.05+00 AW327895.1 EST_HUMAN 24029 37330 7.46 0.05+00 AW327895.1 EST_HUMAN 24029 37330 7.46 0.05+00 AW327895.1 EST_HUMAN 25134 37348 1.89 0.05+00 AW22776.1 EST_HUMAN	Exon NO: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Acession Value Top Hit Acession Source Top Hit Acession Source 24018 37321 4.02 0.0E+00 BE206846.1 EST_HUMAN 24026 37322 4.02 0.0E+00 BE206846.1 EST_HUMAN 24026 37324 3.0E 0.0E+00 BE206846.1 EST_HUMAN 24026 37324 3.8 0.0E+00 AW753028.1 EST_HUMAN 24026 37330 7.46 0.0E+00 AM753028.1 EST_HUMAN 24028 37330 7.46 0.0E+00 AW329577.1 EST_HUMAN 25134 37346 1.89 0.0E+00 AW3292776.1 EST_HUMAN 23173 36401 2.2 0.0E+00 AW282776.1 EST_HUMAN	Exon NO: ORF SEQ ID NO: Expression Signal (Top) Hit Top Hit Acession Source Nalue Top Hit Acession Source Nalue Top Hit Acession Source Source Nalue Top Hit Acession Source Sourc	Exon NO: CAF SEQ ID ID NO: Expression Signal (Top) Hit Top Hit Acession Source Nalue Top Hit Acession Source Source Source Nalue Top Hit Acession Source	Exon NO: CNF SEQ ID ID NO: Signal Signal Most Similar (Top) Hit Top Hit Acession Source 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Cope of the control	Exon NO:- ORF SEQ Expression NO:- Cop Hit Acession Signal Most Similar Top Hit Acession No:- Top Hit Acession Detabase Source So	Exon NO: ORF SEQ ID NO: Expression Signal Most Similar (Top) Hit Pushes Top Hit Acession Pushes Top Hit Acession Source Top Hit Acession Source Top Hit Acession Source 24018 37321 4.02 0.0E+00 BE206846.1 EST HUMAN 24020 37324 4.02 0.0E+00 BE206846.1 EST HUMAN 24020 37324 3.88 0.0E+00 AV55028.1 EST HUMAN 24020 37324 3.89 0.0E+00 AV55028.1 EST HUMAN 24026 37329 7.46 0.0E+00 AV5502776.1 EST HUMAN 25173 36401 2.2 0.0E+00 AV550577.1 EST HUMAN 23976 3726 1.74 0.0E+00 AV56050.2 EST HUMAN 23977 37294 1.29 0.0E+00 BE186509.2 EST HUMAN 23978 37294 1.29 0.0E+00 BE186509.2 EST HUMAN 23979 37294 1.29 0.0E+00 BE186509.2 EST HUMAN 24003 37324	Exm NO: ORF SEQ ID NO: Expression Signal (Top) Hit Value Top Hit Acession Value Top Hit Acession PLAST E Value Top Hit Acession No: Top Hit Acession Source 24018 37322 4.02 0.0E+00 BE206846.1 EST_HUMAN 24026 37324 3.88 0.0E+00 BR206846.1 EST_HUMAN 24026 37324 3.88 0.0E+00 AM733028.1 EST_HUMAN 24026 37326 3.06 0.0E+00 AM733028.1 EST_HUMAN 24026 373276 1.73 0.0E+00 AM732786.1 EST_HUMAN 253773 37278 1.74 0.0E+00 AM732786.1 EST_HUMAN 23978 37279 1.74 0.0E+00 BE364050.2 EST_HUMAN 23978 37279 1.74 0.0E+00 BE364050.2 EST_HUMAN 23978 37279 1.74 0.0E+00 BE364050.1 EST_HUMAN 23978 37284 7.81 0.0E+00 BE364050.1 EST_HUMAN 24050 37365	Exon NO: ORF SEQ ID NO: Expression Signal (Top) Hit Value Top Hit Acession Value Top Hit Acession Value Top Hit Source 24018 37321 4,02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 4,02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 4,02 0.0E+00 BMX5302.1 EST_HUMAN 24020 37324 3.06 0.0E+00 AM753028.1 EST_HUMAN 24020 37324 3.06 0.0E+00 AM753028.1 EST_HUMAN 23976 37236 7.46 0.0E+00 AM258707.1 EST_HUMAN 23978 37276 1.74 0.0E+00 BE568500.2 EST_HUMAN 23978 37276 1.74 0.0E+00 BE568500.2 EST_HUMAN 23978 37280 1.29 0.0E+00 BE568500.2 EST_HUMAN 23978 37284 1.34 0.0E+00 BE568500.2 EST_HUMAN 24050 37384 1.34 0.0E+00 AA76	Exon NO: ORF SEQ ID NO: Expression Signal (Top) Hit Value Top Hit Acession Value Top Hit Acession Source Top Hit Source 24018 37321 4.02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 4.02 0.0E+00 BE206846.1 EST_HUMAN 24020 37324 4.02 0.0E+00 BMS2088.1 EST_HUMAN 24020 37324 3.06 0.0E+00 AMS53028.1 EST_HUMAN 24020 37324 3.06 0.0E+00 AMS68707.1 EST_HUMAN 24020 37326 7.46 0.0E+00 AMS68707.1 EST_HUMAN 24020 37326 7.46 0.0E+00 AMS68707.1 EST_HUMAN 23976 37286 1.74 0.0E+00 BE266809.2 EST_HUMAN 23976 37286 1.77 0.0E+00 BE566809.2 EST_HUMAN 23976 37286 1.78 0.0E+00 BE566809.2 EST_HUMAN 24005 37281 1.78 0.0E+00 AA7

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Single Exon Probes Explessed in Drain	DORF SEQ Expression (Top) Hit Acession Database Signal BLASTE No. Signal Value Surce	37404 1.92 0.0E+00 BE622317.1 EST_HUMAN	37428 1.42 0.0E+00 AIB39634.1 EST_HUMAN	37434 13.79 0.0E+00 BE748899.1 EST_HUMAN	37435 13.79 0.0E+00 BE748899.1 EST_HUMAN	37447 1.81 0.0E+00 AU141882.1 [EST_HUMAN	37448 1.81 0.0E+00 AU141882.1 EST_HUMAN	37451 2.08 0.0E+00 AW006022.1 EST_HUMAN		37455 3.49 0.0E+00 BF002333.1 ES I_HUMAN	37485 2.88 0.0E+00 AW387776.1 EST_HUMAN	37486 2.88 0.0E+00 AW387776.1 EST_HUMAN	2.41 0.0E+00 AW863777.1 EST_HUMAN	37521 4.76 0.0E+00 11435244 NT	37522 4.76 0.0E+00 11435244 NT	37529 5.87 0.0E+00 U36253.1 NT	37533 2.29 0.0E+00 BE379254.1 [EST_HUMAN	37534 2.29 0.0E+00 BE379254.1 EST_HUMAN	37553 2.22 0.0E+00 BE794758.1 [EST_HUMAN	37554 45.09 0.0E+00 BE879633.1 EST_HUMAN	37560 1.62 0.0E+00 4758827 NT	37561 1.62 0.0E+00 4758827 NT	37565 1.65 0.0E+00 AF053543.1 NT	1.56 0.0E+00 AL163204.2 NT	37572 14.06 0.0E+00 BE40893.1 [EST_HUMAN	37573 1.46 0.0E+00 BE148650.1 EST_HUMAN	37574 2.69 0.0E+00 AF223391.1 NT	37575 2.69 0.0E+00/AF223391.1 NT	30878 1.29 0.0E+00[D26535.1 (NT	30879 1.29 0.0E+00 D26535.1 NT	37576 5.6 0.0E+00 BF681641.1 EST_HUMAN	
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	Probe SEQ ID NO:	11491	1151	11529	11529	11539	11539	11542		11546	11571	11571	11582	11601	11601	11608	11612	11612	11632	11634	11640	11640	11644	11646	11653	11654	11655	11655	11657	11657	11658	

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Single Exon Probes Expressed in Brain	ession (Top) Hit Acession Database Top Hit Descriptor Source Source	1.83 0.0E+00 6006002 NT Homo sapiens glutamate receptor, fonotropic, N-methyl D-escartate 24 (GRIN/24) mBNA	+00 AF272663.1 NT	EST HUMAN	EST HUMAN	0.0E+00 X51755.1 NT	ΓN	0.0E+00 BF309120.1 EST_HUMAN	0.0E+00 BE297175.1 EST HUMAN	HUMAN	1.3 0.0E+00 BE744311.1 [EST HUMAN 601576525F1 NIH MGC 9 Homo spalens CDNA clone INA GE-3R37222 F1	EST_HUMAN	EST HUMAN	EST HUMAN	0.0E+00 AW749184.1 EST HUMAN	0.0E+00 AW749184.1 EST HUMAN	0.0E+00 AW367811.1 EST_HUMAN	0.0E+00 AW367811.1 EST HUMAN	0.0E+00 AU117974.1 EST HUMAN	2.46 0.0E+00/AU117974.1 [EST HUMAN AU117974 HEMBA1 Home sablens cDNA clane HEMBA1002812.5]	-00 U36264.1 NT	0.0E+00 U07223.1 NT	6.64 0.0E+00/Z31708.1 NT H.saplens GLAST1 gene for gillial glutamate transporter, exon6	1838/02.x1 NCI_CGAP_GC8 Home sapiens cDNA clone IMAGE:2243087 3' similar to SW:CG2G_HUMAN	0.0E+00 AL132394 1 FST HIMAN	0.0E+00 BE312542.1 EST HIMAN	0.0E+00 A 190993.1 EST HUMAN	0.0E+00/AB011389.1 NT	-00 AL163246.2 NT	11417862 NT	3.05 0.0E+00 5802973 NT Homo saplens antioxidant protein 1 (AOP1), nuclear gene encoding mitochandrial protein mRNA	-00 AF240786.1 NT	0.0E+00 AL041831.1 EST_HUMAN
	milar Hit T E	0.0E+00	0.0E+00 AF2	0.0E+00 AU1	0.0E+00 BE9	0.0E+00 X51	0.0E+00 X51	0.0E+00 BF3	0.0E+00 BE2	0.0E+00 BE7	0.0E+00 BE7	0.0E+00 BE2	0.0E+00 BE2	0.0E+00 BE2	0.0E+00 AW7	0.0E+00 AW7	0.0E+00 AW3	0.0E+00 AW3	0.0E+00 AU1	0.0E+00/AU1	0.0E+00 U362	0.0E+00 U072	0.0E+00 Z317		0.0E+00 AU13	0.0E+00 BE31	0.0E+00 A/190	0.0E+00/AB01	0.0E+00 AL18	0.0E+00	0.0E+00	0.0E+00 AF24	0.0E+00 AL04
	Expression Signal	1.83	1.5	1.71	1.35	2.5	2.5	15.74	11.96	1.3	1.3	1.43	1.43	1.69	1.68	1.68	2.23	2.23	2.46	2.46	1.31	2.45	6.54	- 36	231	2.27	3.89	1.6	4.9	4.1	3.05	2.59	5.39
	ORF SEQ ID NO:						37605			37671	37672	37678	37679	37705	37710	37711	37713	37714	37720	37721		31859	37741	37760	37762	30800	-	-				30816	
	Exan SEQ ID NO:	24258	17906	24262	24265	24282	24282	25136	24330	24342	24342	24348	24348	24375	24380	24380	24382	24382	24387	24387	14946	18891	24407	24419	24421	25399	26257	24468	24482	24488	24501	25218	26226
	Probe SEQ ID NO:	11662	11684	11667	11870	11687	11687	11728	11737	11751	11751	11757	11757	11785	11790	11790	11792	11792	11797	11797	11808	11821	11822	11835	11837	11879	11893	11902	11921	11929	11947	11982	11893

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	_	_								
Top Hit Descriptor	Homo saplens low density lipoprotein-related protein 2 (LRP2). mRNA	Homo sapiens CST gene for cerebroside sulfotransferase, exon 1, 2, 3, 4, 5	Homo sapiens cleavage and polyadenylation specific factor 1, 160kD subunit (CPSF1), mRNA	Homo saplens chromosome 21 segment HS21C046	Homo saplens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA	Wu93c07.x1 NCI_CGAP_Kld3 Homo saplens cDNA clone IMAGE:2527596 3' similar to TR:Q12844 Q12844 BREAKPOINT CI LISTER REGION PROTEIN CONTINUE TABLES TABLES CONTINUED TO TABLES TABLES CONTINUED TO TABLES TABLES CONTINUED TO TABLES TABLES CONTINUED TO TABLES TABLES CONTINUED TO TABLES CONTI	Homo sapiens chromosome 12 open reading frame 3 (C120RF3) mRNA	Homo saplens dihydropyridine receptor alpha 2 subunit (CACNA2D1) gene, exon 8
Top Hit Database Source	N.	F	Ę	Z	Ę	Ę	LN LN	FST HUMAN	L	L'N
Top Hit Acession No.	6806918	AB029900.1	9558724		6806918	11417862	7657020		3844	00 AF083824.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00			0.0E+00
Expression Signal	3.05	212	2.06	2.66	2.77	1.5	4	1.76	1.37	1.39
ORF SEQ ID NO:	30592		31009		26021	30972		30967	26550	
Exen SEQ ID NO:	17905	24905	24927	25410	13390	24999	25004	25042	13890	25261
Probe SEQ ID NO:	12576	12682	12622	12648	12654	12729	12735	12790	12808	12818
	Exon ORF SEQ Expression (Top) Hit Top Hit Acession Database No. Signal Value Source	Exon No: ORF SEQ ID NO: Expression Signal Most Similar (Top) Hit Acession Signal Top Hit Acession No: Source Source Source Top Hit Acession Signal Top Hit Acession Source Source 17905 30592 3.05 0.0E+00 6806918 NT	Exan No: ORF SEQ ID ID NO: Expression Signal 17905 Most Similar (Top) Hit Acession Signal 17905 Top Hit Acession Source Nalue Source 3.05 Top Hit Acession No: 0.0E+00 AB029900.1 Top Hit Acession Source Source 3.05	Exan No: ORF SEQ ID ID NO: Expression Signal (Top) Hit Acession Signal Most Similar (Top) Hit Acession Signal Top Hit Acession Database No: Top Hit Acession Source Source Source 17905 30592 3.05 0.0E+00 6806918 NT 24905 2.12 0.0E+00 AB029900.1 NT 24927 31009 2.06 0.0E+00 AB0229724 NT	Exon SEQ ID ID NO: ORF SEQ Signal ID NO: Expression Signal Signal Signal Most Similar Palue Value Value Top Hit Acession No. Top Hit Acession Source Source Source 17905 30592 3.05 0.06+00 0.06+00 6806918 0.06+00 NT 24905 2.12 0.06+00 0.06+00 AB029900.1 0.06+00 NT 24927 31009 2.06 0.06+00 0.06+00 AB02946.2 0.06+00 NT	Exon SEQ ID ID NO: ORF SEQ Signal Expression PLAST E Value Most Similar No. Top Hit Acession PLAST E Value Top Hit Acession No. Top Hit Acession Source Source 17905 30592 3.05 0.06+00 6806918 NT 24905 2.12 0.06+00 AB029900.1 NT 24927 31009 2.06 0.06+00 AB029900.1 NT 28410 2.06 0.06+00 AB029900.1 NT 285410 2.06 0.06+00 AB02946.2 NT 13390 26021 2.77 0.06+00 AB029418 NT	Exon SEQ ID ID NO: ORF SEQ Signal ID NO: Expression Signal Signa	Exon SEQ ID ID NO: ORF SEQ Signal Expression PLAST E Value Most Similar No. Top Hit Acession PLAST E No. Top Hit Acession No. Top Hit Acession No. Top Hit Acession Source Source 17905 30592 3.05 0.06+00 8606918 NT 24905 2.06 0.06+00 9558724 NT 24910 2.06 0.06+00 9558724 NT 13390 2.05 0.06+00 9558724 NT 24999 30972 1.5 0.06+00 141477862 NT 24999 30972 1.5 0.06+00 7657020 NT 7657020 NT	Exam Seq 1D NO: ORF SEQ Signal S	Exon Seq 1D NO: ORF SEQ Signal S

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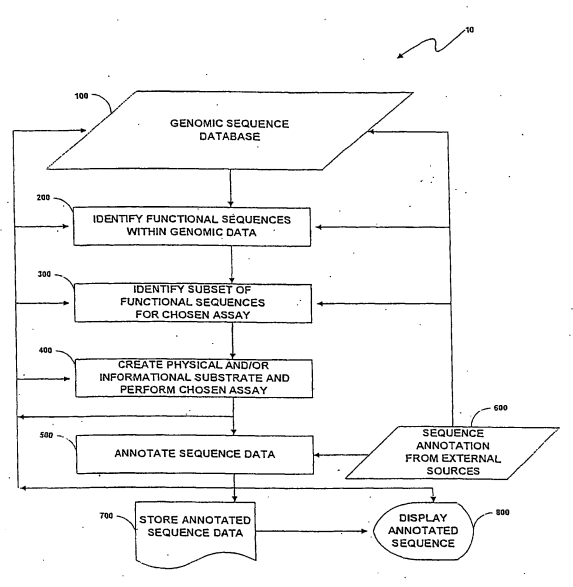


Fig. 1

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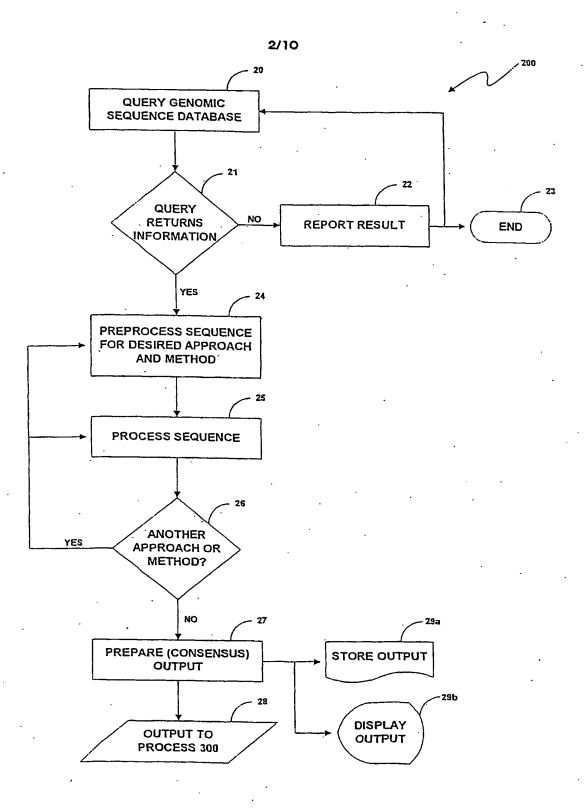


Fig. 2

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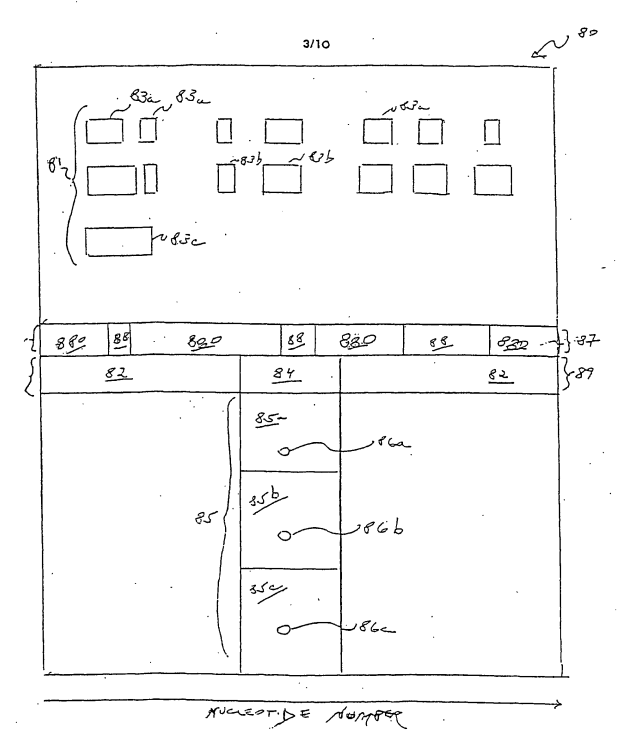


Fig. 3

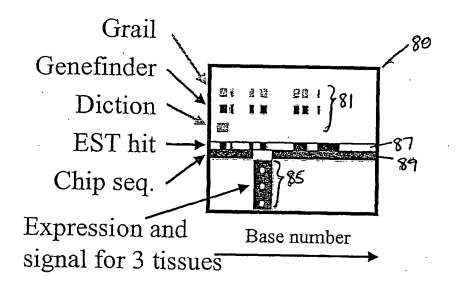


Fig. 4

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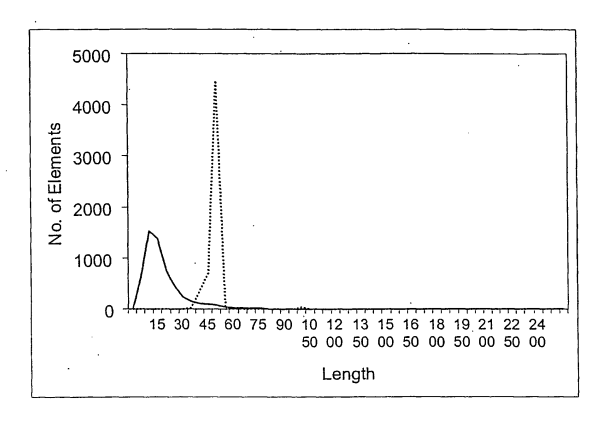


Fig. 5

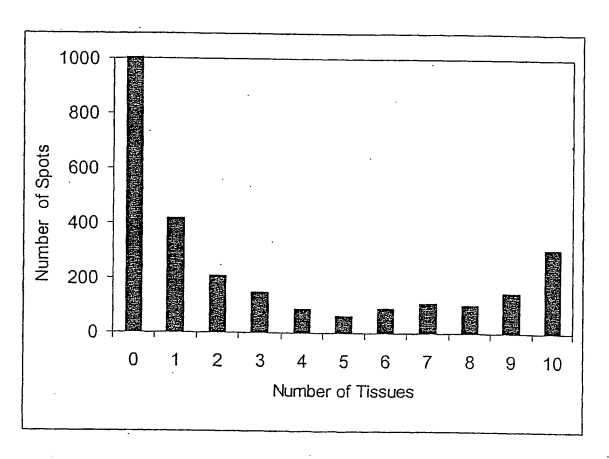
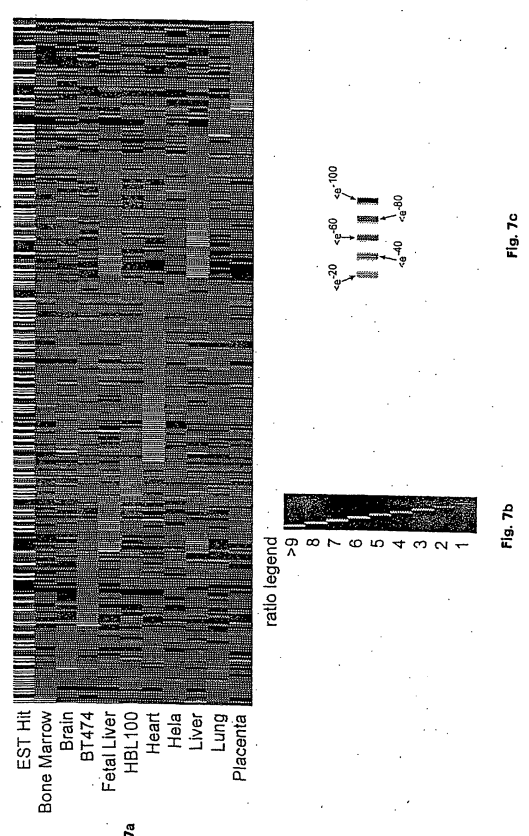


Fig. 6

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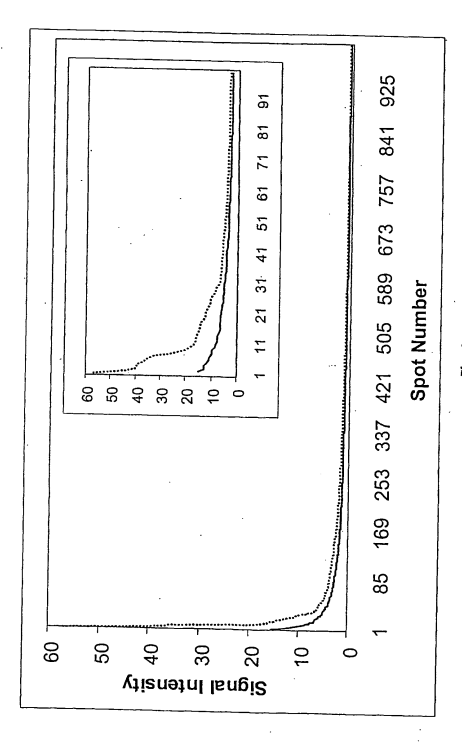


Fig. 8

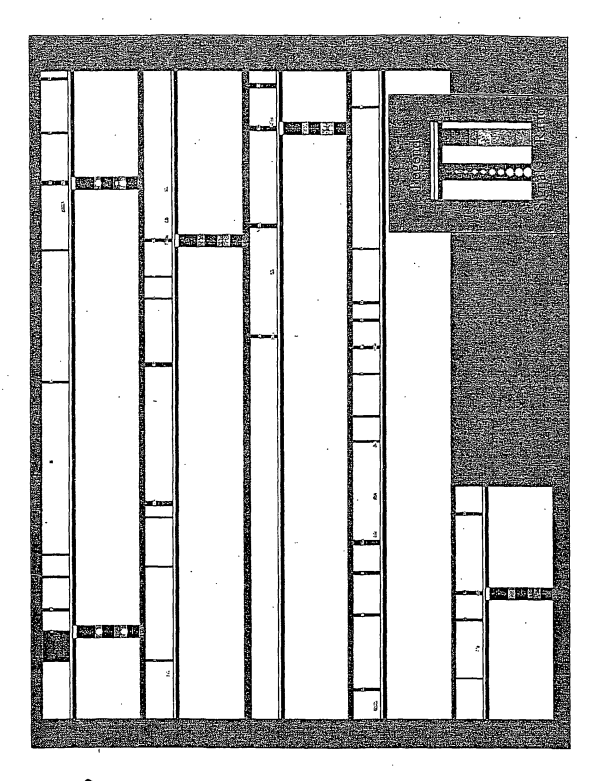


Fig. 9

Fig. 10

